ANNEX 6 Additional Information

6.1. Global Warming Potential Values

Global Warming Potential (GWP) is intended as a quantified measure of the globally averaged relative radiative forcing impacts of a particular greenhouse gas. It is defined as the cumulative radiative forcing–both direct and indirect effects—integrated over a specific period of time from the emission of a unit mass of gas relative to some reference gas (IPCC 2007). Carbon dioxide (CO₂) was chosen as this reference gas. Direct effects occur when the gas itself is a greenhouse gas. Indirect radiative forcing occurs when chemical transformations involving the original gas produce a gas or gases that are greenhouse gases, or when a gas influences other radiatively important processes such as the atmospheric lifetimes of other gases. The relationship between kilotons (kt) of a gas and million metric tons of CO_2 equivalents (MMT CO_2 Eq.) can be expressed as follows:

MMT CO₂ Eq. = (kt of gas) × (GWP) ×
$$\left(\frac{MMT}{1,000 \text{ kt}}\right)$$

where,

MMT $CO_2 Eq.$	=	Million metric tons of CO ₂ equivalent
kt	=	kilotons (equivalent to a thousand metric tons)
GWP	=	Global warming potential
MMT	=	Million metric tons

GWP values allow policy makers to compare the impacts of emissions and reductions of different gases. According to the IPCC, GWP values typically have an uncertainty of ± 35 percent, though some GWP values have larger uncertainty than others, especially those in which lifetimes have not yet been ascertained. In the following decision, the parties to the United Nations Framework Convention on Climate Change (UNFCCC) have agreed to use consistent GWP values from the *IPCC Fourth Assessment Report* (AR4), based upon a 100 year time horizon, although other time horizon values are available (see Table A-267). While this Inventory uses agreed-upon GWP values according to the specific reporting requirements of the UNFCCC, described below, unweighted gas emissions and sinks in kilotons (kt) are provided in the Trends chapter of this report (Table 2-2) and users of the Inventory can apply different metrics and different time horizons to compare the impacts of different greenhouse gases.

...the global warming potential values used by Parties included in Annex I to the Convention (Annex I Parties) to calculate the carbon dioxide equivalence of anthropogenic emissions by sources and removals by sinks of greenhouse gases shall be those listed in the column entitled "Global warming potential for given time horizon" in table 2.14 of the errata to the contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change, based on the effects of greenhouse gases over a 100-year time horizon...¹⁵³

Greenhouse gases with relatively long atmospheric lifetimes (e.g., CO_2 , CH_4 , N_2O , HFCs, PFCs, SF₆, and NF₃) tend to be evenly distributed throughout the atmosphere, and consequently global average concentrations can be determined. However, short-lived gases such as water vapor, carbon monoxide, tropospheric ozone, other indirect greenhouse gases (e.g., NO_x and NMVOCs), and tropospheric aerosols (e.g., SO_2 products and black carbon) vary spatially, and consequently it is difficult to quantify their global radiative forcing impacts. GWP values are generally not attributed to these gases that are short-lived and spatially inhomogeneous in the atmosphere.

¹⁵³ United Nations Framework Convention on Climate Change; http://unfccc.int/resource/docs/2013/cop19/eng/10a03.pdf; 31 January 2014; Report of the Conference of the Parties at its nineteenth session; held in Warsaw from 11 to 23 November 2013; Addendum; Part two: Action taken by the Conference of the Parties at its nineteenth session; Decision 24/CP.19; Revision of the UNFCCC reporting guidelines on annual inventories for Parties included in Annex I to the Convention; p. 2. (UNFCCC 2014)

Table A-267: IPCC AR4 Global Warming Potentials (GWP) and Atmospheric Lifetimes (Years) of Gases	Used in this Report
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Gas	Atmospheric Lifetime	100-year GWP ^a	20-year GWP	500-year GWP
Carbon dioxide (CO ₂)	See footnote ^b	1	1	1
Methane (CH ₄) ^c	12 ^d	25	72	7.6
Nitrous oxide (N ₂ O)	114 ^d	298	289	153
HFC-23	270	14,800	12,000	12,200
HFC-32	4.9	675	2,330	205
HFC-125	29	3,500	6,350	1,100
HFC-134a	14	1,430	3,830	435
HFC-143a	52	4,470	5,890	1,590
HFC-152a	1.4	124	437	38
HFC-227ea	34.2	3,220	5,310	1,040
HFC-236fa	240	9,810	8,100	7,660
HFC-43-10mee	15.9	1,640	4,140	500
CF ₄	50,000 ^d	7,390	5,210	11,200
C_2F_6	10,000	12,200	8,630	18,200
C ₃ F ₈	2,600	8,830	6,310	12,500
C4F10	2,600	8,860	6,330	12,500
c-C ₄ F ₈	3,200	10,300	7,310	14,700
C ₅ F ₁₂	4,100	9,160	6,510	13,300
C ₆ F ₁₄	3,200	9,300	6,600	13,300
SF ₆	3,200	22,800	16,300	32,600
NF3	740	17,200	12,300	20,700

^a GWP values used in this report are calculated over 100 year time horizon.

^b For a given amount of CO₂ emitted, some fraction of the atmospheric increase in concentration is quickly absorbed by the oceans and terrestrial vegetation, some fraction of the atmospheric increase will only slowly decrease over a number of years, and a small portion of the increase will remain for many centuries or more.

• The methane GWP includes the direct effects and those indirect effects due to the production of tropospheric ozone and stratospheric water vapor. The indirect effect due to the production of CO₂ is not included.

^d Methane and N₂O have chemical feedback systems that can alter the length of the atmospheric response, in these cases, global mean atmospheric lifetime (LT) is given first, followed by perturbation time (PT), but only the perturbation time is listed here and not the atmospheric residence time. Source: IPCC (2007)

Table A-268 presents direct GWP values for ozone depleting substances (ODSs). Ozone depleting substances directly absorb infrared radiation and contribute to positive radiative forcing; however, their effect as ozone-depleters also leads to a negative radiative forcing because ozone itself is a potent greenhouse gas. There is considerable uncertainty regarding this indirect effect; direct GWP values are shown, but AR4 does provide a range of net GWP values for ozone depleting substances. The IPCC Guidelines and the UNFCCC do not include reporting instructions for estimating emissions of ODSs because their use is being phased out under the Montreal Protocol (see note below Table A-268). The effects of these compounds on radiative forcing are not addressed in this report.

Table A-268: 100-year Direct Global Warming Potentials for Select Ozone Depleting Substances

Gas	Direct GWP
CFC-11	4,750
CFC-12	10,900
CFC-113	6,130
HCFC-22	1,810
HCFC-123	77
HCFC-124	609
HCFC-141b	725
HCFC-142b	2,310
CH ₃ CCI ₃	146
CCl ₄	1,400
CH₃Br	5
Halon-1211	1,890
Halon-1301	7.140

Note: Because these compounds have been shown to deplete stratospheric ozone, they are typically referred to as ODSs. However, they are also potent greenhouse gases. Recognizing the harmful effects of these compounds on the ozone layer, in 1987 many governments signed the *Montreal Protocol on Substances that Deplete the Ozone Layer* to limit the production and importation of a number of CFCs and other halogenated compounds. The United

States furthered its commitment to phase-out ODSs by signing and ratifying the Copenhagen Amendments to the Montreal Protocol in 1992. Under these amendments, the United States committed to ending the production and importation of halons by 1994, and CFCs by 1996. Source: IPCC (2007)

The IPCC published its Fifth Assessment Report (AR5) in 2013, providing the most current and comprehensive scientific assessment of climate change (IPCC 2013). Within this report, the GWP values were revised relative to the IPCC's Fourth Assessment Report (AR4) (IPCC 2007). Although the AR4 GWP values are used throughout this Inventory report in line with UNFCCC inventory reporting guidelines, it is informative to review the changes to the 100-year GWP values and the impact they have on the total GWP-weighted emissions of the United States. All GWP values use CO₂ as a reference gas; a change in the radiative efficiency of CO₂ thus impacts the GWP of all other greenhouse gases. Since the Second Assessment Report (SAR) and Third Assessment Report (TAR), the IPCC has applied an improved calculation of CO2 radiative forcing and an improved CO₂ response function. The GWP values are drawn from IPCC (2007), with updates for those cases where new laboratory or radiative transfer results have been published. Additionally, the atmospheric lifetimes of some gases have been recalculated, and updated background concentrations were used. Table A-269 shows how the GWP values of the other gases relative to CO_2 tend to be larger in AR4 and AR5 because the revised radiative forcing of CO_2 is lower than in earlier assessments, taking into account revisions in lifetimes. Comparisons of GWP values are based on the 100-year time horizon required for UNFCCC inventory reporting. However, there were some instances in which other variables, such as the radiative efficiency or the chemical lifetime, were altered that resulted in further increases or decreases in particular GWP values in AR5. In addition, the values for radiative forcing and lifetimes have been calculated for a variety of halocarbons. Updates in some well-mixed HFC compounds (including HFC-23, HFC-32, HFC-134a, and HFC-227ea) for AR4 result from investigation into radiative efficiencies in these compounds, with some GWP values changing by up to 40 percent; with this change, the uncertainties associated with these well-mixed HFCs are thought to be approximately 12 percent.

It should be noted that the use of IPCC AR4 GWP values for the current Inventory applies across the entire time series of the Inventory (i.e., from 1990 to 2016). As such, GWP comparisons throughout this chapter are presented relative to AR4 GWPs.

	Lif	etime (years)			GWP (10	0 year)		Difference in GWP (Relative to AR4)					
					•	•	AR5 with				•	AR5 with	AR5 with
Gas	SAR	AR4	AR5	SAR	AR4	AR5 ^a	feedbacks ^b	SAR	SAR (%)	AR5 ^a	AR5 (%)	feedbacks ^b	feedbacks ^b (%)
Carbon dioxide (CO ₂)	C	d	d	1	1	1	1	NC	NC	NC	NC	NC	NC
Methane (CH ₄) ^e	12±3	8.7/12 ^f	12.4	21	25	28	34	(4)	(16%)	3	12%	9	36%
Nitrous oxide (N₂O)	120	120/114 ^f	121	310	298	265	298	12	4%	(33)	(11%)	0	0%
Hydrofluorocarbons													
HFC-23	264	270	222	11,700	14,800	12,400	13,856	(3,100)	(21%)	(2,400)	(16%)	(944)	(6)%
HFC-32	5.6	4.9	5.2	650	675	677	817	(25)	(4%)	2	+%	142	21%
HFC-125	32.6	29	28.2	2,800	3,500	3,170	3,691	(700)	(20%)	(330)	(9%)	191	5%
HFC-134a	14.6	14	13.4	1,300	1,430	1,300	1,549	(130)	(9%)	(130)	(9%)	119	8%
HFC-143a	48.3	52	47.1	3,800	4,470	4,800	5,508	(670)	(15%)	330	7%	1,038	23%
HFC-152a	1.5	1.4	1.5	140	124	138	167	16	13%	14	11%	43	35%
HFC-227ea	36.5	34.2	38.9	2,900	3,220	3,350	3,860	(320)	(10%)	130	4%	640	20%
HFC-236fa	209	240	242	6,300	9,810	8,060	8,998	(3,510)	(36%)	(1,750)	(18%)	(812)	(8)%
HFC-245fa	NA	7.6	7.7	NA	1,030	858	1032	NA	NA	(172)	(17%)	2	+%
HFC-365mfc	NA	6.6	8.7	NA	794	804	966	NA	NA	10	1%	172	22%
HFC-43-10mee	17.1	15.9	16.1	1,300	1,640	1,650	1,952	(340)	(21%)	10	1%	312	19%
Fully Fluorinated Species													
SF ₆	3,200	3,200	3,200	23,900	22,800	23,500	26,087	1,100	5%	700	3%	3,287	14%
CF ₄	50,000	50,000	50,000	6,500	7,390	6,630	7,349	(890)	(12%)	(760)	(10%)	(41)	(1)%
C ₂ F ₆	10,000	10,000	10,000	9,200	12,200	11,100	12,340	(3,000)	(25%)	(1,100)	(9%)	140	1%
C ₃ F ₈	2,600	2,600	2,600	7,000	8,830	8,900	9,878	(1,830)	(21%)	70	1%	1,048	12%
C_4F_{10}	2,600	2,600	2,600	7,000	8,860	9,200	10,213	(1,860)	(21%)	340	4%	1,353	15%
c-C4F8	3,200	3,200	3,200	8,700	10,300	9,540	10,592	(1,600)	(16%)	(760)	(7%)	292	3%
C ₅ F ₁₂	4,100	4,100	4,100	7,500	9,160	8,550	9,484	(1,660)	(18%)	(610)	(7%)	324	4%
C ₆ F ₁₄	3,200	3,200	3,100	7,400	9,300	7,910	8,780	(1,900)	(20%)	(1,390)	(15%)	(520)	(6)%
NF ₃	NA	740	500	NA	17,200	16,100	17,885	NA	NA	(1,100)	(6%)	685	4%

Table A-269: Comparison of GWP values and Lifetimes Used in the SAR, AR4, and AR5

+ Does not exceed 0.05 or 0.05 percent.

NC (No Change)

NA (Not Applicable)

^a The GWP values presented here are the ones most consistent with the methodology used in the AR4 report.

^b The GWP values presented here from the AR5 report include climate-carbon feedbacks for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

• For a given amount of CO₂ emitted, some fraction of the atmospheric increase in concentration is quickly absorbed by the oceans and terrestrial vegetation, some fraction of the atmospheric increase will only slowly decrease over a number of years, and a small portion of the increase will remain for many centuries or more.

^d No single lifetime can be determined for CO₂ (see IPCC 2007).

e The methane GWP includes the direct effects and those indirect effects due to the production of tropospheric ozone and stratospheric water vapor. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

^f Methane and N₂O have chemical feedback systems that can alter the length of the atmospheric response, in these cases, global mean residence time is given first, followed by perturbation time. Note: Parentheses indicate negative values. Source: IPCC (2013), IPCC (2007), IPCC (1996). The choice of GWP values between the SAR, AR4, and AR5 with or without climate-carbon feedbacks has an impact on both the overall emissions estimated by the Inventory, as well as the trend in emissions over time. To summarize, Table A-270 shows the overall trend in U.S. greenhouse gas emissions, by gas, from 1990 through 2016 using the four GWP sets. The table also presents the impact of SAR and AR5 GWP values with or without feedbacks on the total emissions for 1990 and for 2016.

Gas	Difference	Revisio	Revisions to Annual Emission Estimates (Relative to AR4)							
					SAR	AR5 ^a	AR5 ^b	SAR	AR5 ^a	AR5 ^b
	SAR	AR4	AR5 ^a	AR5 ^b		1990			2016	
CO ₂	189.6	189.6	189.6	189.6	NC	NC	NC	NC	NC	NC
CH ₄	(102.9)	(122.5)	(137.2)	(166.6)	(124.8)	93.6	280.8	(105.2)	78.9	236.7
N ₂ O	15.4	14.8	13.2	14.8	14.3	(39.3)	NC	14.9	(40.9)	NC
HFCs, PFCs, SF _{6,}										
and NF ₃	62.6	73.8	73.1	90.5	(11.9)	(9.0)	1.3	(23.2)	(9.7)	17.9
Total	164.6	155.7	138.6	128.2	(122.4)	45.3	282.0	(113.5)	28.3	254.6
Percent Change	2.6%	2.4%	2.2%	1.9%	-1.9%	0.7%	4.4%	-1.7%	0.4%	3.9%

	Table A	-270:	Effects on U.S.	Greenhouse	Gas Emissi	ons Using SA	R. AR4. and	AR5 GWP va	lues (MMT (CO2 Ea.)
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NC (No Change)

^a The GWP values presented here are the ones most consistent with the methodology used in the AR4 report.

^b The GWP values presented here from the AR5 report include climate-carbon feedbacks for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

Note: Totals may not sum due to independent rounding. Excludes sinks. Parentheses indicate negative values.

When the GWP values from the SAR are applied to the emission estimates presented in this report, total emissions for the year 2016 are 6,397.8 MMT CO_2 Eq., as compared to the official emission estimate of 6,511.3 MMT CO_2 Eq. using AR4 GWP values (i.e., the use of SAR GWPs results in a 1.7 percent decrease relative to emissions estimated using AR4 GWPs). Table A-271 provides a detailed summary of U.S. greenhouse gas emissions and sinks for 1990 through 2016, using the GWP values from the SAR. The percent change in emissions for a given gas resulting from using different GWPs is equal to the percent change in the GWP; however, in cases where emissions of multiple gases are combined, as with HFCs or PFCs, the percent change will be a function of the relative quantity of the individual gases. Table A-272 summarizes the resulting change in emissions from using SAR GWP values relative to emissions using AR4 values for 1990 through 2016, including the percent change for 2016.

Table A-971	Recent Trends in II S	Greenhouse Gas Emissio	ns and Sinks using the	SAR GWP values (MMT CO ₂ Fn)
1 auiu #-271.		טו ככווווטעפכ עמפ בוווופפוט	ווס מווע סוווגס עסוווץ נווט	

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Gas/Source	1990	2005	2012	2013	2014	2015	2016
CO ₂	5,121.3	6,132.0	5,366.7	5,519.6	5,568.8	5,420.8	5,310.9
Fossil Fuel Combustion	4,740.3	5,746.9	5,024.4	5,156.9	5,200.3	5,049.3	4,966.0
Electric Power	1,820.8	2,400.9	2,022.2	2,038.1	2,038.0	1,900.7	1,809.3
Transportation	1,467.6	1,855.8	1,661.9	1,677.6	1,717.1	1,735.5	1,782.6
Industrial	858.8	855.7	812.9	843.3	824.9	809.5	809.1
Residential	338.3	357.8	282.5	329.7	345.3	316.8	292.5
Commercial	227.2	227.0	201.3	225.7	233.6	245.4	231.3
U.S. Territories	27.6	49.7	43.5	42.5	41.4	41.4	41.4
Non-Energy Use of Fuels	119.5	138.9	108.0	123.5	118.9	125.6	112.2
Iron and Steel Production &							
Metallurgical Coke Production	101.6	68.2	55.6	53.5	58.4	47.8	42.3
Cement Production	33.5	46.2	35.3	36.4	39.4	39.9	39.4
Petrochemical Production	21.2	26.8	26.5	26.4	26.5	28.1	28.1
Natural Gas Systems	29.8	22.5	23.3	24.8	25.3	24.9	25.5
Petroleum Systems	7.7	11.7	19.3	22.6	26.3	28.8	22.8
Lime Production	11.7	14.6	13.8	14.0	14.2	13.3	12.9
Ammonia Production	13.0	9.2	9.4	10.0	9.6	10.9	12.2
Other Process Uses of Carbonates	6.3	7.6	9.1	11.5	13.0	12.3	11.0
Incineration of Waste	8.0	12.5	10.4	10.4	10.6	10.7	10.7
Urea Fertilization	2.4	3.5	4.3	4.4	4.5	4.9	5.1
Carbon Dioxide Consumption	1.5	1.4	4.0	4.2	4.5	4.5	4.5
Urea Consumption for Non-Agricultural							
Purposes	3.8	3.7	4.4	4.1	1.5	4.2	4.0

Liming	4.7	4.3	6.0	3.9	3.6	3.8	3.9
Ferroalloy Production	2.2	1.4	1.9	1.8	1.9	2.0	1.8
Soda Ash Production	1.4	1.7	1.7	1.7	1.7	1.7	1.7
Titanium Dioxide Production	1.2	1.8	1.5	1.7	1.7	1.6	1.6
Aluminum Production	6.8	4.1	3.4	3.3	2.8	2.8	1.3
Glass Production	1.5	1.9	1.2	1.3	1.3	1.3	1.2
Phosphoric Acid Production	1.5	1.3	1.1	1.1	1.0	1.0	1.0
Zinc Production	0.6	1.0	1.5	1.4	1.0	0.9	0.9
Lead Production Silicon Carbida Braduction and	0.5	0.0	0.5	0.5	0.5	0.5	0.5
Consumption	0.4	0.2	0.2	0.2	0.2	0.2	0.2
Consumption	0.4	0.2	0.2	0.2	0.2	0.2	0.2
Abandoned Oil and Gas Wells	+	+	+	+	+	+	+
Magnesium Production and Processing	+	+	+	+	+	+	+
Wood Biomass, Ethanol, and Biodiesel	040.4	000 7	007 7	040.4	004.0	040.4	000.0
Consumption ^a	219.4	230.7	287.7	316.4	324.3	310.4	309.3
	103.0	113.1 570 A	100.8 EEG E	99.8 EEC E	103.4 667 7	110.9 550 0	110.0
Entoric Formontation	137.0	3/0.4	000.0 1/0 1	330.3 130.0	337.7 137.0	130.0	332.2
Natural Gas Systems	163.9	141.0	140.1	135.0	138.0	139.9	142.9
l andfille	150.8	142.1	08.3	95.1	Q/ 7	03.8	۹۵ <i>۱</i>
Manure Management	31.2	47.3	55.1	53.1	52.8	55.0	56.9
Coal Mining	81.1	53.9	55.8	54.3	54.2	51.4	45.2
Petroleum Systems	33.4	27.0	27.4	30.7	32.4	32.0	32.4
Wastewater Treatment	13.2	13.3	12.7	12.5	12.6	12.7	12.5
Rice Cultivation	13.5	14.0	9.5	9.7	10.7	10.3	11.5
Stationary Combustion	7.2	6.6	6.2	7.4	7.5	6.7	6.2
Abandoned Oil and Gas Wells	5.5	5.8	5.9	5.9	5.9	6.0	6.0
Abandoned Underground Coal Mines	6.0	5.5	5.2	5.2	5.3	5.4	5.6
Mobile Combustion	10.7	7.9	4.3	3.9	3.6	3.2	3.1
Composting	0.3	1.6	1.6	1.7	1.8	1.8	1.8
Field Burning of Agricultural Residues	0.2	0.2	0.2	0.2	0.2	0.2	0.2
Petrochemical Production	0.2	0.1	0.1	0.1	0.1	0.2	0.2
Ferroalloy Production	+	+	+	+	+	+	+
Silicon Carbide Production and							
Consumption	+	+	+	+	+	+	+
Iron and Steel Production &							
Metallurgical Coke Production	+	+	+	+	+	+	+
Incineration of Waste	+	+	+	+	+	+	+
International Bunker Fuels ^o	0.7	0.1	0.1	0.1	0.1	204.0	204.4
Agricultural Soil Management	309.0 260.5	31Z.Z	349.3 257.0	3//.9 287 7	3/3./ 285.0	394.9	304.4 205.0
Stationary Compustion	200.5	203.7	257.9	207.7	205.0	18.8	290.0
Mobile Compustion	11.5	10.2	25.2	23.4	21.5	20.1	10.0
Manure Management	14.6	40.4	18.2	23.4 18.2	18.2	18.4	18.0
Nitric Acid Production	12.6	11.8	10.2	11.1	11.2	12.4	10.5
Adipic Acid Production	15.8	7.4	5.8	4 1	57	4.4	7.3
Wastewater Treatment	3.5	4.6	4.8	4.9	5.0	5.0	5.2
N ₂ O from Product Uses	4.4	4.4	4.4	4.4	4.4	4.4	4.4
Caprolactam, Glyoxal, and Glyoxylic							
Acid Production	1.7	2.2	2.1	2.1	2.1	2.1	2.1
Composting	0.4	1.7	1.8	1.9	1.9	2.0	2.0
Incineration of Waste	0.5	0.4	0.3	0.3	0.3	0.3	0.3
Semiconductor Manufacture	+	0.1	0.2	0.2	0.2	0.2	0.2
Field Burning of Agricultural Residues	0.1	0.1	0.1	0.1	0.1	0.1	0.1
International Bunker Fuels ^b	0.9	1.0	1.0	0.9	0.9	1.0	1.0
HFCs	36.9	107.8	131.0	131.1	135.6	139.0	140.2
Substitution of Ozone Depleting							
Substances ^d	0.3	91.8	126.5	127.7	131.3	135.3	137.6

HCFC-22 Production	36.4	15.8	4.3	3.2	4.0	3.4	2.2
Semiconductor Manufacture	0.2	0.2	0.2	0.1	0.2	0.3	0.3
Magnesium Production and Processing	0.0	0.0	+	0.1	0.1	0.1	0.1
PFCs	20.6	5.6	4.9	4.8	4.7	4.2	3.6
Semiconductor Manufacture	2.2	2.6	2.4	2.3	2.5	2.5	2.4
Aluminum Production	18.4	3.0	2.5	2.5	2.1	1.7	1.1
Substitution of Ozone Depleting							
Substances	0.0	+	+	+	+	+	+
SF ₆	30.2	12.3	7.0	6.6	6.7	6.2	6.5
Electrical Transmission and Distribution	24.2	8.7	4.9	4.7	4.9	4.5	4.5
Magnesium Production and Processing	5.4	2.9	1.7	1.5	1.0	0.9	1.1
Semiconductor Manufacture	0.5	0.7	0.4	0.4	0.8	0.8	0.9
NF ₃	NA						
Semiconductor Manufacture	NA						
Total	6,233.2	7,208.3	6,415.4	6,596.5	6,649.2	6,524.0	6,397.8
LULUCF Emissions ^c	9.7	21.3	24.1	17.8	18.2	35.2	35.1
LULUCF CH ₄ Emissions	5.6	11.1	12.6	9.2	9.4	18.8	18.8
LULUCF N ₂ O Emissions	4.1	10.1	11.5	8.6	8.8	16.4	16.3
LULUCF Carbon Stock Change ^e	(830.2)	(754.2)	(779.5)	(755.0)	(760.0)	(733.4)	(754.9)
LULUCF Sector Net Total ^f	(820.5)	(732.9)	(755.4)	(737.2)	(741.8)	(698.1)	(719.7)
Net Emissions (Sources and Sinks)	5,412.7	6,475.4	5,660.0	5,859.4	5,907.4	5,825.9	5,678.1

Notes: Total emissions presented without LULUCF. Net emissions presented with LULUCF.

+ Does not exceed 0.05 MMT CO₂ Eq.

NA (Not Applicable)

a Emissions from Wood Biomass and Biofuel Consumption are not included specifically in summing energy sector totals. Net carbon fluxes from changes in biogenic carbon reservoirs are accounted for in the estimates for LULUCF.

^b Emissions from International Bunker Fuels are not included in totals.

c LULUCF emissions of CH₄ and N₂O are reported separately from gross emissions totals. LULUCF emissions include the CH₄ and N₂O emissions reported for *Peatlands Remaining Peatlands*, Forest Fires, Drained Organic Soils, Grassland Fires, and *Coastal Wetlands Remaining Coastal Wetlands*; CH₄ emissions from *Land Converted to Coastal Wetlands*; and N₂O emissions from Forest Soils and Settlement Soils.

^d Small amounts of PFC emissions also result from this source.

^e LULUCF Carbon Stock Change is the net C stock change from the following categories: Forest Land Remaining Forest Land, Land Converted to Forest Land, Cropland Remaining Cropland, Land Converted to Cropland, Grassland Remaining Grassland, Land Converted to Grassland, Wetlands Remaining Wetlands, Land Converted to Wetlands, Settlements Remaining Settlements, and Land Converted to Settlements.

^f The LULUCF Sector Net Total is the net sum of all CH₄ and N₂O emissions to the atmosphere plus net carbon stock changes.

Notes: Totals may not sum due to independent rounding. Parentheses indicate net sequestration.

Table A-272: Change in U.S. Greenhouse Gas Emissions Using SAR GWP values relative to AR4 GWP values (MMT CO2 Eq.)

								Percent
Gas/Source	1990	2005	2012	2013	2014	2015	2016	Change in 2016
CO ₂	NC	NC	 NC	NC	NC	NC	NC	NC
CH₄	(124.8)	(110.2)	(106.0)	(106.0)	(106.2)	(106.5)	(105.2)	(16%)
Enteric Fermentation	(26.3)	(27.0)	(26.7)	(26.5)	(26.3)	(26.6)	(27.2)	(16%)
Natural Gas Systems	(31.2)	(27.1)	(25.5)	(26.2)	(26.3)	(26.6)	(26.2)	(16%)
Landfills	(28.7)	(21.2)	(18.7)	(18.1)	(18.0)	(17.9)	(17.2)	(16%)
Manure Management	(5.9)	(9.0)	(10.5)	(10.1)	(10.1)	(10.6)	(10.8)	(16%)
Coal Mining	(15.4)	(10.3)	(10.6)	(10.3)	(10.3)	(9.8)	(8.6)	(16%)
Petroleum Systems	(6.4)	(5.1)	(5.2)	(5.9)	(6.2)	(6.1)	(6.2)	(16%)
Wastewater Treatment	(2.5)	(2.5)	(2.4)	(2.4)	(2.4)	(2.4)	(2.4)	(16%)
Rice Cultivation	(2.6)	(2.7)	(1.8)	(1.8)	(2.0)	(2.0)	(2.2)	(16%)
Stationary Combustion	(1.4)	(1.3)	(1.2)	(1.4)	(1.4)	(1.3)	(1.2)	(16%)
Abandoned Oil and Gas Wells	(1.0)	(1.1)	(1.1)	(1.1)	(1.1)	(1.1)	(1.1)	(16%)
Abandoned Underground Coal Mines	(1.2)	(1.1)	(1.0)	(1.0)	(1.0)	(1.0)	(1.1)	(16%)
Mobile Combustion	(2.0)	(1.5)	(0.8)	(0.8)	(0.7)	(0.6)	(0.6)	(16%)
Composting	(0.1)	(0.3)	(0.3)	(0.3)	(0.3)	(0.3)	(0.3)	(16%)
Field Burning of Agricultural								
Residues	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
Petrochemical Production	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)

Ferroalloy Production	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
Silicon Carbide Production and	()		()	()	()		()	(100()
Consumption	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
Iron and Steel Production &				()	()		()	(400())
Metallurgical Coke Production	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
Incineration of Waste	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
International Bunker Fuels ^a	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(16%)
N2U	14.3	14.4	13.5	14.0	14.5	15.3	14.9	4%
Agricultural Soll Management	10.1	10.2	10.0	11.1	11.0	11.9	11.4	4%
Stationary Compustion	0.4	0.7	0.7	0.8	0.8	0.7	0.7	4%
Mobile Combustion	1.7	1.6	1.0	0.9	0.8	0.8	0.7	4%
Manure Management	0.6	0.7	0.7	0.7	0.7	0.7	0.7	4%
Nitric Acid Production	0.5	0.5	0.4	0.4	0.4	0.5	0.4	4%
Adipic Acid Production	0.6	0.3	0.2	0.2	0.2	0.2	0.3	4%
Wastewater Treatment	0.1	0.2	0.2	0.2	0.2	0.2	0.2	4%
N ₂ O from Product Uses	0.2	0.2	0.2	0.2	0.2	0.2	0.2	4%
Caprolactam, Glyoxal, and Glyoxylic								
Acid Production	0.1	0.1	0.1	0.1	0.1	0.1	0.1	4%
Composting	+	0.1	0.1	0.1	0.1	0.1	0.1	4%
Incineration of Waste	+	+	+	+	+	+	+	4%
Semiconductor Manufacture	+	+	+	+	+	+	+	4%
Field Burning of Agricultural	_							
Residues	+	+	+	+	+	+	+	
International Bunker Fuels ^a	+	+	+	+	+	+	+	4%
HFCs	(9.7)	(15.2)	(19.6)	(20.0)	(21.1)	(21.8)	(22.1)	(14%)
Substitution of Ozone Depleting	_							
Substances ^b	+	(10.9)	(18.4)	(19.1)	(20.0)	(20.8)	(21.5)	(13%)
HCFC-22 Production	(9.7)	(4.2)	(1.1)	(0.9)	(1.1)	(0.9)	(0.6)	(21%)
Semiconductor Manufacture	(+)	(+)	(+)	(+)	(0.1)	(0.1)	(0.1)	(21%)
Magnesium Production and	_							
Processing	0.0	0.0	(+)	(+)	(+)	(+)	(+)	(9%)
PFCs	(3.6)	(1.1)	(1.0)	(1.0)	(0.9)	(0.9)	(0.7)	(17%)
Semiconductor Manufacture	(+)	(0.7)	(0.6)	(0.5)	(0.6)	(0.6)	(0.5)	(18%)
Aluminum Production	(3.0)	(0.5)	(0.4)	(0.4)	(0.4)	(0.3)	(0.2)	(16%)
Substitution of Ozone Depleting	_							
Substances	0.0	(+)	(+)	(+)	(+)	(+)	(+)	(12%)
SF ₆	1.4	0.6	0.3	0.3	0.3	0.3	0.3	5%
Electrical Transmission and	_							
Distribution	1.1	0.4	0.2	0.2	0.2	0.2	0.2	5%
Magnesium Production and	_							
Processing	0.3	0.1	0.1	0.1	+	+	+	5%
Semiconductor Manufacture	+	+	+	+	+	+	+	5%
NF ₃	NA							
Semiconductor Manufacture	NA							
Total	(122.4)	(112.0)	(113.4)	(112.6)	(113.9)	(114.1)	(113.5)	(1.7%)

NC (No Change)

NA (Not Applicable)

+ Absolute value does not exceed 0.05 MMT CO₂ Eq. ^a Emissions from International Bunker Fuels are not included in totals.

Emissions from international Bunker Fuels are not included in totals
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^b Small amounts of PFC emissions also result from this source.

Notes: Totals may not sum due to independent rounding. Parentheses indicate negative values.

Table A-273 below shows a comparison of total emissions estimates by sector using both the IPCC SAR and AR4 GWP values. For most sectors, the change in emissions that result from using SAR relative to AR4 GWP values was minimal. The effect on emissions from waste was by far the greatest (15.0 percent decrease in 2016 using SAR GWP values, relative to emissions using AR4 GWP values), due the predominance of CH_4 emissions in this sector. Emissions from all other sectors were comprised of mainly CO_2 or a mix of gases, which moderated the effect of the changes.

Sector	1990	_	2005	2012	2013	2014	2015	2016
Energy								
AR4 GWP, Used In Inventory	5,325.1		6,285.2	5,511.2	5,671.4	5,715.4	5,567.8	5,455.2
SAR GWP	5,268.6		6,240.1	5,467.3	5,626.4	5,670.0	5,522.8	5,411.8
Difference (%)	(1.1%)		(0.7%)	(0.8%)	(0.8%)	(0.8%)	(0.8%)	(0.8%)
Industrial Processes and								
Product Use								
AR4 GWP, Used In Inventory	342.0		358.6	357.4	357.9	371.4	367.8	362.1
SAR GWP	331.4		343.4	337.5	337.6	350.1	345.8	339.8
Difference (%)	(3.1%)		(4.2%)	(5.6%)	(5.7%)	(5.7%)	(6.0%)	(6.1%)
Agriculture								
AR4 GWP, Used In Inventory	489.2		520.0	519.8	543.1	539.8	566.9	562.6
SAR GWP	465.0		492.2	491.4	516.5	513.2	540.2	534.5
Difference (%)	(4.9%)		(5.4%)	(5.5%)	(4.9%)	(4.9%)	(4.7%)	(5.0%)
LULUCF								
AR4 GWP, Used In Inventory	(819.6)		(731.1)	(753.5)	(735.8)	(740.4)	(695.2)	(716.8)
SAR GWP	(820.5)		(732.9)	(755.4)	(737.2)	(741.8)	(698.1)	(719.7)
Difference (%)	0.1%		0.2%	0.3%	0.2%	0.2%	0.4%	0.4%
Waste								
AR4 GWP, Used In Inventory	199.3		156.4	140.4	136.7	136.5	135.6	131.5
SAR GWP	168.2		132.6	119.2	116.1	116.0	115.3	111.8
Difference (%)	(15.6%)		(15.2%)	(15.1%)	(15.0%)	(15.0%)	(15.0%)	(15.0%)
Net Emissions (Sources and								
Sinks)								
AR4 GWP, Used In Inventory	5,536.0		6,589.1	5,775.3	5,973.3	6,022.8	5,942.9	5,794.5
SAR GWP	5,412.7		6,475.4	5,660.0	5,859.4	5,907.4	5,825.9	5,678.1
Difference (%)	(2.2%)		(1.7%)	(2.0%)	(1.9%)	(1.9%)	(2.0%)	(2.0%)

Table A-273: Comparison of Emissions by Sector using IPCC AR4 and SAR GWP Values (MMT CO2Eq.)

Notes: Totals may not sum due to independent rounding. Parentheses indicate negative values.

Further, Table A-274 and Table A-275 show the comparison of emission estimates using AR5 GWP values relative to AR4 GWP values without climate-carbon feedbacks for the non-CO₂ gases, on an emissions and percent change basis. Table A-276 and Table A-277 show the comparison of emission estimates using AR5 GWP values with climate-carbon feedbacks. The use of AR5 GWP values without climate-carbon feedbacks¹⁵⁴ results in an increase in emissions of CH₄ and SF₆ relative to AR4 GWP values, but a decrease in emissions of other gases. The use of AR5 GWP values with climatecarbon feedbacks does not impact CO₂ and N₂O emissions; however, it results in an increase in emissions of CH₄, SF₆, and NF₃ relative to AR4 GWP values, and has mixed impacts on emissions of other gases. Overall, these comparisons of AR4 and AR5 GWP values do not have a significant effect on calculated U.S. emissions, resulting in an increase in emissions of less than 1 percent using AR5 GWP values, or approximately 4 percent when using AR5 GWP values with climate-carbon feedbacks. As with the comparison of SAR and AR4 GWP values presented above, the percent change in emissions is equal to the percent change in the GWP for each gas; however, in cases where multiple gases are emitted in varying amounts the percent change is variable over the years, such as with Substitution of Ozone Depleting Substances.

¹⁵⁴ The IPCC AR5 report provides additional information on emission metrics. See https://www.ipcc.ch/pdf/assessment-report/ar5/wg1/WG1AR5_Chapter08_FINAL.pdf.

Gas	1990	2005	2012	2013	2014	2015	2016
CO ₂	NC						
CH ₄	93.6	82.6	79.5	79.5	79.7	79.8	78.9
N ₂ O	(39.3)	(39.6)	(37.2)	(40.2)	(40.0)	(42.0)	(40.9)
HFCs	(7.5)	(10.6)	(9.4)	(9.0)	(9.3)	(9.5)	(9.5)
PFCs	(2.4)	(0.6)	(0.6)	(0.6)	(0.5)	(0.5)	(0.4)
SF ₆	0.9	0.4	0.2	0.2	0.2	0.2	0.2
NF ₃	(+)	(+)	(+)	(+)	(+)	(+)	(+)
Total	45.3	32.1	32.5	29.9	30.0	27.9	28.3

Table A-274: Change in U.S. Greenhouse Gas Emissions Using AR5ª without Climate-Carbon Feedbacks Relative to AR4 GWP Values (MMT CO2 Eq.)

+ Absolute value does not exceed 0.05 MMT CO2 Eq.

NC (No Change)

^a The GWP values presented here are the ones most consistent with the methodology used in the AR4 report. The AR5 report has also calculated GWP values (shown in Table A-276) where climate-carbon feedbacks have been included for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

Notes: Total emissions presented without LULUCF. Totals may not sum due to independent rounding. Parentheses indicate negative values.

Table A-275: Change in U.S. Greenhouse Gas Emissions Using AR5 ^a without Climate-Carbon Feedback	s Relative to AR4 GWP
Values (Percent)	

Gas/Source	1990	2005	2012	2013	2014	2015	2016
CO ₂	NC						
CH4	12.0%	12.0%	12.0%	12.0%	12.0%	12.0%	12.0%
N ₂ O	(11.1%)	(11.1%)	(11.1%)	(11.1%)	(11.1%)	(11.1%)	(11.1%)
SF ₆	3.1%	3.1%	3.1%	3.1%	3.1%	3.1%	3.1%
NF ₃	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)
HFCs	(16.0%)	(8.6%)	(6.2%)	(6.0%)	(6.0%)	(5.9%)	(5.8%)
Substitution of Ozone	. ,			. ,			. ,
Depleting Substances	11.3%	(7.1%)	(5.8%)	(5.7%)	(5.6%)	(5.6%)	(5.6%)
HCFC-22 Production ^b	(16.2%)	(16.2%)	(16.2%)	(16.2%)	(16.2%)	(16.2%)	(16.2%)
Semiconductor Manufacture ^c	(16.2%)	(16.2%)	(16.2%)	(16.2%)	(16.2%)	(16.1%)	(16.2%)
Magnesium Production and	. ,		. ,	. ,	. ,	. ,	. ,
Processing ^d	0.0%	0.0%	(9.1%)	(9.1%)	(9.1%)	(9.1%)	(9.1%)
PFCs	(10.0%)	(9.6%)	(9.5%)	(9.6%)	(9.5%)	(9.5%)	(9.4%)
Semiconductor Manufacture ^c	(9.4%)	(9.2%)	(9.1%)	(9.2%)	(9.2%)	(9.2%)	(9.2%)
Aluminum Production ^e	(10.1%)	(10.1%)	(10.0%)	(10.0%)	(10.0%)	(10.0%)	(9.9%)
Substitution of Ozone	. ,		. ,	. ,	. ,	. ,	. ,
Depleting Substances ^{d,f}	0.0%	(10.3%)	(10.3%)	(10.3%)	(10.3%)	(10.3%)	(10.3%)
Total	0.7%	0.4%	0.5%	0.4%	0.4%	0.4%	0.4%

NC (No Change)

^a The GWP values presented here are the ones most consistent with the methodology used in the AR4 report. The AR5 report has also calculated GWP values (shown in Table A-277) where climate-carbon feedbacks have been included for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

^b HFC-23 emitted.

^c Emissions from HFC-23, CF₄, C₂F₆, C₃F₈, C₄F₈, SF₆, as well as other HFCs and PFCs used as heat transfer fluids.

^d Zero change in beginning of time series since emissions were zero.

e PFC emissions from CF4 and C2F6.

^f PFC emissions from CF₄.

Note: Total emissions presented without LULUCF. Parentheses indicate negative values.

Gas	1990	2005	2012	2013	2014	2015	2016
CO ₂	NC						
CH ₄	280.8	247.9	238.5	238.5	239.0	239.5	236.7
N ₂ O	NC						
HFCs	(2.9)	8.9	15.2	15.7	16.1	16.6	17.0
PFCs	(+)	+	+	+	+	+	+
SF ₆	4.2	1.7	1.0	0.9	0.9	0.9	0.9
NF ₃	+	+	+	+	+	+	+
Total	282.0	258.5	254.7	255.1	256.1	257.1	254.6

Table A-276: Change in U.S. Greenhouse Gas Emissions Using AR5 with Climate-Carbon Feedbacks^a Relative to AR4 GWP Values (MMT CO₂ Eq.)

+ Absolute value does not exceed 0.05 MMT CO₂ Eq.

NC (No Change)

^a The GWP values presented here from the AR5 report include climate-carbon feedbacks for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

Notes: Total emissions presented without LULUCF. Totals may not sum due to independent rounding. Parentheses indicate negative values.

Table A-277:	Change in U.S.	Greenhouse Gas	Emissions Using	NR5 with Climate	-Carbon Feedbacks ^a	Relative to A	R4 GWP
Values (Perc	ent)						

Gas/Source	1990	2005	2012	2013	2014	2015	2016
CO ₂	NC	NC	NC	NC	NC	NC	NC
CH ₄	36.0%	36.0%	36.0%	36.0%	36.0%	36.0%	36.0%
N ₂ O	NC	NC	NC	NC	NC	NC	NC
SF ₆	14.4%	14.4%	14.4%	14.4%	14.4%	14.4%	14.4%
NF ₃	4.0%	4.0%	4.0%	4.0%	4.0%	4.0%	4.0%
HFCs	(6.1%)	7.2%	10.1%	10.4%	10.3%	10.3%	10.5%
Substitution of Ozone							
Depleting Substances	34.7%	9.9%	10.8%	10.9%	10.9%	10.8%	10.8%
HCFC-22 Production ^b	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)	(6.4%)
Semiconductor	, ,			, ,	ι, γ	、 <i>,</i>	、
Manufacture ^c	(6.4%)	(6.3%)	(6.3%)	(6.3%)	(6.3%)	(6.3%)	(6.3%)
Magnesium Production and							
Processing ^d	0.0%	0.0%	8.3%	8.3%	8.3%	8.3%	8.3%
PFCs	(0.2%)	0.3%	0.4%	0.3%	0.4%	0.4%	0.5%
Semiconductor							
Manufacture ^c	0.6%	0.9%	0.9%	0.9%	0.8%	0.8%	0.7%
Aluminum Production ^e	(0.3%)	(0.3%)	(0.2%)	(0.2%)	(0.1%)	(0.1%)	+%
Substitution of Ozone							
Depleting Substances ^{d,f}	0.0%	(0.6%)	(0.6%)	(0.6%)	(0.6%)	(0.6%)	(0.6%)
Total	4.4%	3.5%	3.9%	3.8%	3.8%	3.9%	3.9%

NC (No Change)

+ Does not exceed 0.05 percent.

^a The GWP values presented here from the AR5 report include climate-carbon feedbacks for the non-CO₂ gases in order to be consistent with the approach used in calculating the CO₂ lifetime. Additionally, the AR5 reported separate values for fossil versus biogenic methane in order to account for the CO₂ oxidation product.

^b HFC-23 emitted.

^c Emissions from HFC-23, CF₄, C₂F₆, C₃F₈, C₄F₈, SF₆, as well as other HFCs and PFCs used as heat transfer fluids.

^d Zero change in beginning of time series since emissions were zero.

^e PFC emissions from CF₄ and C₂F₆.

^f PFC emissions from CF₄.

Notes: Total emissions presented without LULUCF. Parentheses indicate negative values. Excludes Sinks.

6.2. Ozone Depleting Substance Emissions

Ozone is present in both the stratosphere,¹⁵⁵ where it shields the earth from harmful levels of ultraviolet radiation, and at lower concentrations in the troposphere,¹⁵⁶ where it is the main component of anthropogenic photochemical "smog." Chlorofluorocarbons (CFCs), halons, carbon tetrachloride, methyl chloroform, and hydrochlorofluorocarbons (HCFCs), along with certain other chlorine and bromine containing compounds, have been found to deplete the ozone levels in the stratosphere. These compounds are commonly referred to as ozone depleting substances (ODSs). If left unchecked, stratospheric ozone depletion could result in a dangerous increase of ultraviolet radiation reaching the earth's surface. In 1987, nations around the world signed the *Montreal Protocol on Substances that Deplete the Ozone Layer*. This landmark agreement created an international framework for limiting, and ultimately eliminating, the production of most ozone depleting substances. ODSs have historically been used in a variety of industrial applications, including refrigeration and air conditioning, foam blowing, fire extinguishing, sterilization, solvent cleaning, and as an aerosol propellant.

In the United States, the Clean Air Act Amendments of 1990 provide the legal instrument for implementation of the *Montreal Protocol* controls. The Clean Air Act classifies ozone depleting substances as either Class I or Class II, depending upon the ozone depletion potential (ODP) of the compound.¹⁵⁷ The production of CFCs, halons, carbon tetrachloride, and methyl chloroform—all Class I substances—has already ended in the United States. However, large amounts of these chemicals remain in existing equipment,¹⁵⁸ and stockpiles of the ODSs, as well as material recovered from equipment being decommissioned, are used for maintaining the existing equipment. As a result, emissions of Class I compounds will continue, albeit generally in decreasing amounts, for many more years. Class II designated substances, all of which are HCFCs, have been, or are being, phased out at later dates than Class I compounds because they have lower ODPs. These compounds served, and in some cases continue to serve, as interim replacements for Class I compounds in many industrial applications. The use and emissions of HCFCs in the United States is anticipated to continue for several decades as equipment that use Class II substances are retired from use. Under current *Montreal Protocol* controls, however, the production for domestic use of all HCFCs in the United States must end by the year 2030.

In addition to contributing to ozone depletion, CFCs, halons, carbon tetrachloride, methyl chloroform, and HCFCs are also potent greenhouse gases. However, the depletion of the ozone layer has a cooling effect on the climate that counteracts the direct warming from tropospheric emissions of ODSs. Stratospheric ozone influences the earth's radiative balance by absorption and emission of longwave radiation from the troposphere as well as absorption of shortwave radiation from the sun; overall, stratospheric ozone has a warming effect.

The IPCC has prepared both direct GWP values and net (combined direct warming and indirect cooling) GWP ranges for some of the most common ozone depleting substances (IPCC 2007). See Annex 6.1, Global Warming Potential Values, for a listing of the direct GWP values for ODS.

Although the IPCC emission inventory guidelines do not require the reporting of emissions of ozone depleting substances, the United States believes that the inventory presents a more complete picture of climate impacts when we include these compounds. Emission estimates for several ozone depleting substances are provided in Table A-278.

Compound	1990	2005	2012	2013	2014	2015	2016
Class I							
CFC-11	29	12	24	24	24	25	25
CFC-12	132	22	5	5	4	4	3
CFC-113	59	17	2	0	0	0	0
CFC-114	4	1	+	+	0	0	0
CFC-115	8	2	+	+	+	+	+

Table A-2/8: Emissions of Uzone Depleting Substa	nces (kt)
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¹⁵⁵ The stratosphere is the layer from the top of the troposphere up to about 50 kilometers. Approximately 90 percent of atmospheric ozone is within the stratosphere. The greatest concentration of ozone occurs in the middle of the stratosphere, in a region commonly called the ozone layer.

¹⁵⁶ The troposphere is the layer from the ground up to about 11 kilometers near the poles and 16 kilometers in equatorial regions (i.e., the lowest layer of the atmosphere, where humans live). It contains roughly 80 percent of the mass of all gases in the atmosphere and is the site for weather processes including most of the water vapor and clouds.

¹⁵⁷ Substances with an ozone depletion potential of 0.2 or greater are designated as Class I. All other designated substances that deplete stratospheric ozone but which have an ODP of less than 0.2 are Class II.

¹⁵⁸ Older refrigeration and air-conditioning equipment, fire extinguishing systems, and foam products blown with CFCs/HCFCs may still contain Class I ODS.

Carbon Tetrachloride	4	0	0	0	0	0	0
Methyl Chloroform	223	0	0	0	0	0	0
Halon-1211	2	1	1	1	1	1	+
Halon-1301	2	+	+	+	+	+	+
Class II							
HCFC-22	49	82	70	67	63	59	54
HCFC-123	0	1	1	1	1	1	1
HCFC-124	0	2	1	1	1	+	+
HCFC-141b	1	4	9	10	10	9	9
HCFC-142b	1	4	1	1	2	2	3
HCFC-225ca/cb	0	+	+	+	+	+	+

+ Does not exceed 0.5 kt.

Methodology and Data Sources

Emissions of ozone depleting substances were estimated using the EPA's Vintaging Model. The model, named for its method of tracking the emissions of annual "vintages" of new equipment that enter into service, is a "bottom-up" model. It models the consumption of chemicals based on estimates of the quantity of equipment or products sold, serviced, and retired each year, and the amount of the chemical required to manufacture and/or maintain the equipment. The Vintaging Model makes use of this market information to build an inventory of the in-use stocks of the equipment in each of the enduses. Emissions are estimated by applying annual leak rates, service emission rates, and disposal emission rates to each population of equipment. By aggregating the emission and consumption output from the different end-uses, the model produces estimates of total annual use and emissions of each chemical. Please see Annex 3.9, Methodology for Estimating HFC and PFC Emissions from Substitution of Ozone Depleting Substances, of this Inventory for a more detailed discussion of the Vintaging Model.

Uncertainties

Uncertainties exist with regard to the levels of chemical production, equipment sales, equipment characteristics, and end-use emissions profiles that are used by these models. Please see the Substitution of Ozone Depleting Substances section of this report for a more detailed description of the uncertainties that exist in the Vintaging Model.

6.3. Sulfur Dioxide Emissions

Sulfur dioxide (SO₂), emitted into the atmosphere through natural and anthropogenic processes, affects the Earth's radiative budget through photochemical transformation into sulfate aerosols that can (1) scatter sunlight back to space, thereby reducing the radiation reaching the Earth's surface; (2) affect cloud formation; and (3) affect atmospheric chemical composition (e.g., stratospheric ozone, by providing surfaces for heterogeneous chemical reactions). The overall effect of SO₂-derived aerosols on radiative forcing is believed to be negative (IPCC 2007). However, because SO₂ is short-lived and unevenly distributed through the atmosphere, its radiative forcing impacts are highly uncertain. Sulfur dioxide emissions have been provided below in Table A-279.

The major source of SO_2 emissions in the United States is the burning of sulfur containing fuels, mainly coal. Metal smelting and other industrial processes also release significant quantities of SO_2 . The largest contributor to U.S. emissions of SO_2 is electricity generation, accounting for 43.8 percent of total SO_2 emissions in 2016 (see Table A-280); coal combustion accounted for approximately 92.0 percent of that total. The second largest source was industrial fuel combustion, which produced 20.2 percent of 2016 SO_2 emissions. Overall, SO_2 emissions in the United States decreased by 88.3 percent from 1990 to 2016. The majority of this decline came from reductions from electricity generation, primarily due to increased consumption of low sulfur coal from surface mines in western states.

Sulfur dioxide is important for reasons other than its effect on radiative forcing. It is a major contributor to the formation of urban smog and acid rain. As a contributor to urban smog, high concentrations of SO_2 can cause significant increases in acute and chronic respiratory diseases. In addition, once SO_2 is emitted, it is chemically transformed in the atmosphere and returns to earth as the primary contributor to acid deposition, or acid rain. Acid rain has been found to accelerate the decay of building materials and paints, cause the acidification of lakes and streams, and damage trees. As a result of these harmful effects, the United States has regulated the emissions of SO_2 under the Clean Air Act. The EPA has also developed a strategy to control these emissions via four programs: (1) the National Ambient Air Quality Standards program,¹⁵⁹ (2) New Source Performance Standards,¹⁶⁰ (3) the New Source Review/Prevention of Significant Deterioration Program,¹⁶¹ and (4) the Sulfur Dioxide Allowance Program.¹⁶²

Sector/Source	1990	2005	2012	2013	2014	2015	2016
Energy	19,628	12,364	5,271	5,270	3,859	2,950	1,959
Stationary Sources	18,407	11,541	5,006	5,005	3,640	2,756	1,790
Oil and Gas Activities	390	180	108	108	93	93	93
Mobile Sources	793	619	142	142	95	70	44
Waste Combustion	38	25	15	15	32	32	32
Industrial Processes and							
Product Use	1,307	831	604	604	496	496	496
Other Industrial Processes	362	327	171	171	156	156	156
Miscellaneous ^a	11	114	179	179	135	135	135
Chemical and Allied Product							
Manufacturing	269	228	115	115	104	104	104
Metals Processing	659	158	131	131	98	98	98
Storage and Transport	6	2	8	8	3	3	3
Solvent Use	0	+	+	+	+	+	+
Degreasing	0	0	0	0	0	0	0
Graphic Arts	0	0	0	0	0	0	0
Dry Cleaning	NA	0	0	0	0	0	0
Surface Coating	0	0	0	0	0	0	0
Other Industrial	0	+	+	+	+	+	+
Nonindustrial	NA	NA	NA	NA	NA	NA	NA
Agriculture	NA	NA	NA	NA	NA	NA	NA
Agricultural Burning	NA	NA	NA	NA	NA	NA	NA
Waste	+	1	+	+	1	1	1

Table A-279: SO₂ Emissions (kt)

¹⁵⁹ [42 U.S.C § 7409, CAA § 109]

¹⁶⁰ [42 U.S.C § 7411, CAA § 111]

¹⁶¹ [42 U.S.C § 7473, CAA § 163]

¹⁶² [42 U.S.C § 7651, CAA § 401]

Landfills + 1 + + 1 1 1 1 Wastewater Treatment + 0									
Landfills + 1 + + 1	Total	20,935	13,196	;	5,876	5,874	4,357	3,448	2,457
Landfills + 1 + + 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 0	Miscellaneous ^a	+	()	0	0	0	0	0
Landfills + 1 1 + + 1 1 1	Wastewater Treatment	+	()	0	0	0	0	0
	Landfills	+			+	+	1	1	1

+ Does not exceed 0.5 kt
NA (Not Applicable)
a Miscellaneous includes other combustion and fugitive dust categories.
Note: Totals may not sum due to independent rounding.
Source: Data taken from EPA (2016) and disaggregated based on EPA (2003).

Table A-280: SO₂ Emissions from Electricity Generation (kt)

Fuel Type	1990	20)5	2012	2013	2014	2015	2016
Coal	13,808	8,6	30	3,858	3,856	2,690	1,877	989
Oil	580	4	58	203	203	142	99	52
Gas	1	1	74	77	77	54	38	20
Internal Combustion	45		57	25	25	18	12	7
Other	NA		71	31	31	22	15	8
Total	14,433	9,4	39	4,195	4,194	2,925	2,041	1,075

NA (Not Applicable) Note: Totals may not sum due to independent rounding. Source: Data taken from EPA (2016) and disaggregated based on EPA (2003).

6.4. Complete List of Source Categories

Chapter/Source	Gas(es)
Energy	
Eossil Fuel Combustion	CO_2
Non-Energy Use of Fossil Fuels	CO_2
Stationary Combustion (excluding CO ₂)	$CH_4 N_2 O CO NO_2 NMVOC$
Mobile Combustion (excluding CO ₂)	CH_4 N ₂ O CO NO _x NMVOC
Coal Mining	CH4
Abandoned Underground Coal Mines	CH4
Petroleum Systems	CH4
Natural Gas Systems	CH4
Abandoned Oil and Gas Wells	
Incineration of Waste	CO_2 CH ₄ N ₂ O NO ₂ CO NMVOC
Industrial Processes and Product Use	
Cement Production	CO ₂
Lime Production	
Glass Production	
Other Process Lises of Carbonates	CO2
Ammonia Production	
Urea Consumption for Non-Agricultural Purposes	
Nitric Acid Production	N ₂ O
Adjuic Acid Production	N ₂ O
Caprolactam Glyoxal and Glyoxylic Production	N ₂ O
Silicon Carbide Production and Consumption	$CO_2 CH_4$
	CO_2
Soda Ash Production	
Petrochemical Production	
HCEC-22 Production	HEC-23
Carbon Dioxide Consumption	CO2
Phosphoric Acid Production	
Iron and Steel Production & Metallurgical Coke Production	
Ferroallov Production	CO_2 CH ₄
Aluminum Production	CO_2 CE4 C2E6
Magnesium Production and Processing	CO_2 HECS SE
Lead Production	CO_2
Zinc Production	
Semiconductor Manufacture	N2O HECS PECS ^a SE6 NE3
Substitution of Ozone Depleting Substances	HECs PECs ^b
Electrical Transmission and Distributing	SE
N2O from Product Uses	N ₂ O
Agriculture	1120
Enteric Fermentation	CH4
Manure Management	CH4 N2O
Rice Cultivation	CH ₄
Liming	CO ₂
Urea Fertilization	CO ₂
Field Burning of Agricultural Residues	CH_4 N ₂ O NO ₂ CO
Agricultural Soil Management	N ₂ O
I and Use, I and Use Change, and Forestry	120
Forest Land Remaining Forest Land	CO2, CH4, N2O, NOx, CO
Land Converted to Forest Land	CO ₂
Cropland Remaining Cropland	
Land Converted to Cropland	CO ₂
Grassland Remaining Grassland	CO_2 CH_4 N_2O_2 NO_2 CO_2
Land Converted to Grassland	CO ₂
Wetlands Remaining Wetlands	CO2. CH4. N2O
Land Converted to Wetlands	CO ₂ , CH ₄

Settlements Remaining Settlements	CO ₂ , N ₂ O
Land Converted to Settlements	CO ₂
Waste	
Landfills	CH4, NO _x , CO, NMVOC
Wastewater Treatment	CH ₄ , N ₂ O, NO _x , CO, NMVOC
Composting	CH4, N2O

 Includes HFC-23, CF₄, C₂F₆, as well as other HFCs and PFCs used as heat transfer fluids.
Includes HFC-23, HFC-32, HFC-125, HFC-134a, HFC-143a, HFC-236fa, CF₄, HFC-152a, HFC-227ea, HFC-245fa, HFC-4310mee, and PFC/PFPEs.

 \circ The LULUCF Sector includes CH₄ and N₂O emissions to the atmosphere and net carbon stock changes. The term "flux" is used to describe the net emissions of greenhouse gases accounting for both the emissions of CO₂ to and the removals of CO₂ from the atmosphere. Removal of CO₂ from the atmosphere is also referred to as "carbon sequestration."

6.5. Constants, Units, and Conversions

Metric Prefixes

Although most activity data for the United States is gathered in customary U.S. units, these units are converted into metric units per international reporting guidelines. Table A-281 provides a guide for determining the magnitude of metric units.

Table A-281: Guide to Metric Unit Prefixes

Prefix/Symbol	Factor
atto (a)	10 ⁻¹⁸
femto (f)	10 ⁻¹⁵
pico (p)	10 ⁻¹²
nano (n)	10 ⁻⁹
micro (µ)	10 ⁻⁶
milli (m)	10 ⁻³
centi (c)	10 ⁻²
deci (d)	10 ⁻¹
deca (da)	10
hecto (h)	10 ²
kilo (k)	10 ³
mega (M)	10 ⁶
giga (G)	10 ⁹
tera (T)	10 ¹²
peta (P)	10 ¹⁵
exa (E)	10 ¹⁸

Unit Conversions

1 kilogram 1 pound 1 short ton 1 metric ton	= = =	2.205 pounds 0.454 kilograms 2,000 pounds 1,000 kilograms	= =	0.9072 metric tons 1.1023 short tons	3	
1 cubic meter 1 cubic foot 1 U.S. gallon 1 barrel (bbl) 1 barrel (bbl) 1 liter	= = = =	35.315 cubic feet 0.02832 cubic meters 3.785412 liters 0.159 cubic meters 42 U.S. gallons 0.001 cubic meters	3			
1 foot 1 meter 1 mile 1 kilometer	= = =	0.3048 meters 3.28 feet 1.609 kilometers 0.622 miles				
1 acre 1 square mile	= =	43,560 square feet 2.589988 square kilo	= mete	0.4047 hectares	s =	4,047 square meters
Degrees Celsius Degrees Kelvin	= =	(Degrees Fahrenheit Degrees Celsius + 27	– 32) 73.15	*5/9		

Density Conversions¹⁶³

Methane	1 cubic meter	=	0.67606 kilogra	ms	
Carbon dioxide	1 cubic meter	=	1.85387 kilogra	ms	
			-		
Natural gas liquids	1 metric ton	=	11.6 barrels	=	1,844.2 liters
Unfinished oils	1 metric ton	=	7.46 barrels	=	1,186.04 liters
Alcohol	1 metric ton	=	7.94 barrels	=	1,262.36 liters
Liquefied petroleum gas	1 metric ton	=	11.6 barrels	=	1,844.2 liters
Aviation gasoline	1 metric ton	=	8.9 barrels	=	1,415.0 liters
Naphtha jet fuel	1 metric ton	=	8.27 barrels	=	1,314.82 liters
Kerosene jet fuel	1 metric ton	=	7.93 barrels	=	1,260.72 liters
Motor gasoline	1 metric ton	=	8.53 barrels	=	1,356.16 liters
Kerosene	1 metric ton	=	7.73 barrels	=	1,228.97 liters
Naphtha	1 metric ton	=	8.22 barrels	=	1,306.87 liters
Distillate	1 metric ton	=	7.46 barrels	=	1,186.04 liters
Residual oil	1 metric ton	=	6.66 barrels	=	1,058.85 liters
Lubricants	1 metric ton	=	7.06 barrels	=	1,122.45 liters
Bitumen	1 metric ton	=	6.06 barrels	=	963.46 liters
Waxes	1 metric ton	=	7.87 barrels	=	1,251.23 liters
Petroleum coke	1 metric ton	=	5.51 barrels	=	876.02 liters
Petrochemical feedstocks	1 metric ton	=	7.46 barrels	=	1,186.04 liters
Special naphtha	1 metric ton	=	8.53 barrels	=	1,356.16 liters
Miscellaneous products	1 metric ton	=	8.00 barrels	=	1,271.90 liters

Energy Conversions

Converting Various Energy Units to Joules

The common energy unit used in international reports of greenhouse gas emissions is the joule. A joule is the energy required to push with a force of one Newton for one meter. A terajoule (TJ) is one trillion (10^{12}) joules. A British thermal unit (Btu, the customary U.S. energy unit) is the quantity of heat required to raise the temperature of one pound of water one degree Fahrenheit at or near 39.2 degrees Fahrenheit.

1 TJ = 2.388×10¹¹ calories 23.88 metric tons of crude oil equivalent 947.8 million Btus 277,800 kilowatt-hours

Converting Various Physical Units to Energy Units

Data on the production and consumption of fuels are first gathered in physical units. These units must be converted to their energy equivalents. The conversion factors in Table A-282 can be used as default factors, if local data are not available. See Appendix A of EIA's *Monthly Energy Review, February 2018* (EIA 2018) for more detailed information on the energy content of various fuels.

¹⁶³ Reference: EIA (2007)

Table A-282:	Conversion	Factors to	Eneray	v Units (He	eat Eo	uivalents)
IUNIU A LUL.	0011101 31011	1 4 4 4 4 4 4 4	LIIUISI		յաւ եպ	urvurvnicos

Fuel Type (Units)	Factor
Solid Fuels (Million Btu/Short ton)	
Anthracite coal	22.573
Bituminous coal	23.89
Sub-bituminous coal	17.14
Lignite	12.866
Coke	23.367
Natural Gas (Btu/Cubic foot)	1,037
Liquid Fuels (Million Btu/Barrel)	
Motor gasoline	5.059
Aviation gasoline	5.048
Kerosene	5.670
Jet fuel, kerosene-type	5.670
Distillate fuel	5.773
Residual oil	6.287
Naphtha for petrochemicals	5.248
Petroleum coke	6.104
Other oil for petrochemicals	5.825
Special naphthas	5.248
Lubricants	6.065
Waxes	5.537
Asphalt	6.636
Still gas	6.287
Misc. products	5.796

Note: For petroleum and natural gas, *Monthly Energy Review, February 2018* (EIA 2018). For coal ranks, *State Energy Data Report 1992* (EIA 1993). All values are given in higher heating values (gross calorific values).

6.6. Abbreviations

ABS	Acrylonitrile butadiene styrene
AC	Air conditioner
ACC	American Chemistry Council
AEDT	FAA Aviation Environmental Design Tool
AEO	Annual Energy Outlook
AF&PA	American Forest and Paper Association
AFEAS	Alternative Fluorocarbon Environmental Acceptability Study
AFOLU	Agriculture, Forestry, and Other Land Use
AFV	Alternative fuel vehicle
AGA	American Gas Association
AHEF	Atmospheric and Health Effect Framework
AHRI	Air-Conditioning, Heating, and Refrigeration Institute
AISI	American Iron and Steel Institute
ANGA	American Natural Gas Alliance
ANL	Argonne National Laboratory
APC	American Plastics Council
API	American Petroleum Institute
APTA	American Public Transportation Association
AR4	IPCC Fourth Assessment Report
AR5	IPCC Fifth Assessment Report
ARI	Advanced Resources International
ARMA	Autoregressive moving-average
ARMS	Agricultural Resource Management Surveys
ASAE	American Society of Agricultural Engineers
ASTM	American Society for Testing and Materials
AZR	American Zinc Recycling
BCEF	Biomass conversion and expansion factors
BEA	Bureau of Economic Analysis, U.S. Department of Commerce
BLM	Bureau of Land Management
BoC	Bureau of Census
BOD	Biological oxygen demand
BOD5	Biochemical oxygen demand over a 5-day period
BOEM	Bureau of Ocean Energy Management
BOEMRE	Bureau of Ocean Energy Management, Regulation and Enforcement
BOF	Basic oxygen furnace
BRS	Biennial Reporting System
BTS	Bureau of Transportation Statistics, U.S. Department of Transportation
Btu	British thermal unit
С	Carbon
C&D	Construction and demolition waste
C&EN	Chemical and Engineering News
CAAA	Clean Air Act Amendments of 1990
CaO	Calcium Oxide
CAPP	Canadian Association of Petroleum Producers
CARB	California Air Resources Board
CBI	Confidential business information
C-CAP	Coastal Change Analysis Program
CDAP	Chemical Data Access Tool
CEAP	USDA-NRCS Conservation Effects Assessment Program
CEFM	Cattle Enteric Fermentation Model
CEMS	Continuous emission monitoring system
CFC	Chlorotluorocarbon
CFR	Lode of Federal Regulations
CGA	Compressed Gas Association
CH4	Metnane
CHP	Combined heat and power

CI	Confidence interval
CIGRE	International Council on Large Electric Systems
CKD	Cement kiln dust
CLE	Crown Light Exposure
CMA	Chemical Manufacturer's Association
CMM	Coal mine methane
CMOP	Coalbed Methane Outreach Program
CMP	Chemical Market Peporter
CNC	
	Compressed natural gas
00	
	Carbon dioxide
COGCC	Colorado Oil and Gas Conservation Commission
CRF	Common Reporting Format
CRM	Component ratio method
CRP	Conservation Reserve Program
CSRA	Carbon Sequestration Rural Appraisals
CTIC	Conservation Technology Information Center
CVD	Chemical vapor deposition
CWNS	Clean Watershed Needs Survey
d.b.h	Diameter breast height
DE	Digestible energy
DESC	Defense Energy Support Center-DoD's defense logistics agency
DFAMS	Defense Fuels Automated Management System
DHS	Department of Homeland Security
DM	Dry matter
DOC	Degradable organic carbon
DOC	U.S. Department of Commerce
	U.S. Department of Defense
	U.S. Department of Energy
DOE	U.S. Department of the Interior
DOI	U.S. Department of Transportation
DUI	
	Direct Reduced Iron
EAF	Electric arc furnace
EDB	Aircraft Engine Emissions Databank
EDF	Environmental Defense Fund
EER	Energy economy ratio
EF	Emission factor
EFMA	European Fertilizer Manufacturers Association
EJ	Exajoule
EGR	Exhaust gas recirculation
EGU	Electric generating unit
EIA	Energy Information Administration, U.S. Department of Energy
EIIP	Emissions Inventory Improvement Program
EOR	Enhanced oil recovery
FPA	U.S. Environmental Protection Agency
FRS	Economic Research Service
EV	Electric vehicle
	Encound Vegetation Index
	Food and Agricultural Organization
FAU	Food and Agricultural Organization database
FAUSIAI	Food and Agricultural Organization database
	Framework Convention on Climate Change
FEB	
FERC	Federal Energy Regulatory Commission
FGD	Flue gas desulfurization
FHWA	Federal Highway Administration
FIA	Forest Inventory and Analysis
FIADB	Forest Inventory and Analysis Database

FIPR	Florida Institute of Phosphate Research
FOD	First order decay
FQSV	First-quarter of silicon volume
FSA	Farm Service Agency
FTP	Federal Test Procedure
a	Gram
GaAs	Gallium Arsenide
GCV	Gross calorific value
CDP	Gross damestic product
GHC	Groophouse gas
CHCPD	Greenhouse gas
CI	
GJ	Gudajoule Cult Offenere Activity Data System
GUADS	
GPG	Good Practice Guidance
GRI	Gas Research Institute
GSAM	Gas Systems Analysis Model
GII	Gas Technology Institute
GWP	Global warming potential
ha	Hectare
HBFC	Hydrobromofluorocarbon
HC	Hydrocarbon
HCFC	Hydrochlorofluorocarbon
HCFO	Hydrochlorofluoroolefin
HDDV	Heavy duty diesel vehicle
HDGV	Heavy duty gas vehicle
HDPE	High density polyethylene
HF	Hydraulically fractured
HFC	Hvdrofluorocarbon
HEO	Hydrofluoroolefin
HFF	Hydrofluoroethers
HHV	Higher Heating Value
ΗΜΔ	Hot Mix Asphalt
	Hospital/medical/infectious waste incinerator
	Host Transfor Eluid
ште	Harmonized Tariff Schodulo
	Integrated Circuit
ICAU	International Civil Aviation Organization
ICE	Internal combustion engine
IEA	International Energy Agency
IFO	Intermediate Fuel Oil
IISRP	International Institute of Synthetic Rubber Products
ILENR	Illinois Department of Energy and Natural Resources
IMO	International Maritime Organization
IPAA	Independent Petroleum Association of America
IPCC	Intergovernmental Panel on Climate Change
IPPU	Industrial Processes and Product Use
ITC	U.S. International Trade Commission
ITRS	International Technology Roadmap for Semiconductors
JWR	Jim Walters Resources
KCA	Key category analysis
ka	Kilogram
kt	Kiloton
kWh	Kilowatt hour
IDDT	Light-duty diesel truck
	Light-duty diesel vehicle
LDGT	Light-duty deservenice
	Light-duty gas index
LDGV	Light-duty gas vehicle

LDPE	Low density polyethylene
LDT	Light-duty truck
LDV	Light-duty vehicle
LEV	Low emission vehicles
LFG	Landfill gas
LFGE	Landfill gas-to-energy
LHV	Lower Heating Value
IKD	Lime kiln dust
	Linear low density polyethylene
IMOP	EPA's Landfill Methane Outreach Program
LNG	Liquefied natural das
L PG	Liquefied netroleum gas(es)
	Landing and take-off
	Land Lise Land-Lise Change and Forestry
	International Convention for the Prevention of Pollution from Shins
MC	Motorovole
MCE	Methane conversion factor
MCI	Maximum Contaminant Levels
MCED	Thousand oubic foot por day
MOL	Metered dese inhalors
MECS	EIA Monufacturaria Energy Consumption Survey
	LiA Manuacturer's Energy Consumption Survey
	Micro-electromechanical systems
	Monuniy Energy Review
Mao	Manne yas oli
MU	Magnesiulio
	Mejajoule Major Land Pasouroo Aroa
mm	Major Land Resource Area Millimeter
	Million Pritich thormal units
	Million cubic foot
	Million cubic feet
MMS	Miniori cubic reel per uay Minerals Management Service
MMT	Million Metric Tons
	Million metric tons carbon equivalent
	Million metric tons carbon diovide equivalent
	Moderate Resolution Imaging Spectroradiometer
Moll	Memorandum of Understanding
MOVES	IIS EDA's Motor Vehicle Emission Simulator model
MPG	Miles per gallon
MRIC	Multi-Resolution Land Characteristics Consortium
MRV	Monitoring reporting and verification
MSHA	Mine Safety and Health Administration
MSW	Municipal solid waste
MT	Metric ton
MTRE	Methyl Tertiary Butyl Ether
MTBS	Monitoring Trends in Burn Severity
MVAC	Motor vehicle air conditioning
MY	Model vear
N ₂ O	Nitrous oxide
NA	Not available
NACWA	National Association of Clean Water Agencies
NAHMS	National Animal Health Monitoring System
NAICS	North American Industry Classification System
NAPAP	National Acid Precipitation and Assessment Program
NARR	North American Regional Reanalysis Product
NAS	National Academies of Sciences, Engineering, and Medicine
NASA	National Aeronautics and Space Administration
NASF	National Association of State Foresters
NASS	USDA's National Agriculture Statistics Service

NC	No change
NCASI	National Council of Air and Stream Improvement
NCV	Net calorific value
NE	Not estimated
NEI	National Emissions Inventory
NEMA	National Electrical Manufacturers Association
NEMS	National Energy Modeling System
NESHAP	National Emission Standards for Hazardous Air Pollutants
NEU	Non-Energy Lise
NEV	Neighborhood Electric Vehicle
NE ₂	Nitrogen trifluoride
NGHGI	National Greenhouse Gas Inventory
NGI	Natural das liquids
NID	National Inventory Doport
	National line Association
	National Line Association
NLCD	National Land Cover Dataset
NMOC	Non-methane organic compounds
NMVOC	Non-methane volatile organic compound
NMOG	Non-methane organic gas
NO	Nitric oxide
NO	Not occurring
NO ₂	Nitrogen Dioxide
NOx	Nitrogen oxides
NOAA	National Oceanic and Atmospheric Administration
NPDES	National Pollutant Discharge Elimination System
NPRA	National Petroleum and Refiners Association
NRC	National Research Council
NRCS	Natural Resources Conservation Service
NRI	National Resources Inventory
NSCEP	National Service Center for Environmental Publications
NSCR	Non-selective catalytic reduction
NSPS	New source performance standards
NWS	National Weather Service
OAG	Official Airline Guide
	EPA Office of Atmospheric Programs
	EPA Office of Air Quality Planning and Standards
	Ozone depleting potential
	Organization of Economic Co operation and Development
OECD	
OGJ	
OH	Hydroxyl radical
OMS	EPA Office of Mobile Sources
ORNL	Oak Ridge National Laboratory
USHA OTA	Occupational Safety and Health Administration
OTA	Office of Technology Assessment
OTAQ	EPA Office of Transportation and Air Quality
PAH	Polycyclic aromatic hydrocarbons
PCC	Precipitate calcium carbonate
PDF	Probability Density Function
PECVD	Plasma enhanced chemical vapor deposition
PET	Polyethylene terephthalate
PET	Potential evapotranspiration
PEVM	PFC Emissions Vintage Model
PFC	Perfluorocarbon
PFPE	Perfluoropolyether
PHMSA	Pipeline and Hazardous Materials Safety Administration
PI	Productivity index
PLS	Pregnant liquor solution

POTW	Publicly Owned Treatment Works
ppbv	Parts per billion (10 ⁹) by volume
ppm	Parts per million
ppmv	Parts per million (10 ⁶) by volume
potv	Parts per trillion (10^{12}) by volume
PRP	Pasture/Range/Paddock
DQ	Polyetyrene
	Drimony Comple Linit
P30	
PU	Polyuretnane
PVC	Polyvinyl chloride
PV	Photovoltaic
QA/QC	Quality Assurance and Quality Control
QBtu	Quadrillion Btu
R&D	Research and Development
RECs	Reduced Emissions Completions
RCRA	Resource Conservation and Recovery Act
RMA	Rubber Manufacturers' Association
RPA	Resources Planning Act
	Pogrossion through the origin
	Cegiession-unough-une-ongin
SAE	Society of Automotive Engineers
SAGE	System for assessing Aviation's Global Emissions
SAN	Styrene Acrylonitrile
SAR	IPCC Second Assessment Report
SCR	Selective catalytic reduction
SCSE	South central and southeastern coastal
SEC	Securities and Exchange Commission
SEMI	Semiconductor Equipment and Materials Industry
SE	Sulfur hexafluoride
SiC	Silicon Carbida
	Somiconductor International Canacity Statistics
	Semiconductor International Capacity Statistics
SNAP	
SNG	Synthetic natural gas
SO_2	Sulfur dioxide
SOC	Soil Organic Carbon
SOG	State of Garbage survey
SOHIO	Standard Oil Company of Ohio
SSURGO	Soil Survey Geographic Database
STMC	Scrap Tire Management Council
SULEV	Super Ultra Low Emissions Vehicle
SWANA	Solid Waste Association of North America
SWDS	Solid waste disposal sites
	Traatad anaarabically (waatawatar)
	Trained entred mean
TAM	Typical animal mass
TAME	l ertiary amyl methyl ether
TAR	IPCC Third Assessment Report
TBtu	Trillion Btu
TDN	Total digestible nutrients
TEDB	Transportation Energy Data Book
TFI	The Fertilizer Institute
TIGER	Topologically Integrated Geographic Encoding and Referencing survey
T.I	Teraioule
TIFV	Traditional low emissions vehicle
	Total Manufactured Laver Area
	Tovia Palagaa Inventary
	I UNIC INCICASE INVENTIONY
ISUF	mazardous waste treatment, storage, and disposal facility
IVA	I ennessee Valley Authority
UAN	Urea ammonium nitrate
UDI	Utility Data Institute
UFORE	U.S. Forest Service's Urban Forest Effects model

UG	Underground (coal mining)
U.S.	United States
U.S. ITC	United States International Trade Commission
UEP	United Egg Producers
ULEV	Ultra low emission vehicle
UNEP	United Nations Environmental Programme
UNFCCC	United Nations Framework Convention on Climate Change
USAA	U.S. Aluminum Association
USAF	United States Air Force
USDA	United States Department of Agriculture
USFS	United States Forest Service
USGS	United States Geological Survey
VAIP	EPA's Voluntary Aluminum Industrial Partnership
VAM	Ventilation air methane
VKT	Vehicle kilometers traveled
VMT	Vehicle miles traveled
VOCs	Volatile organic compounds
VS	Volatile solids
WBJ	Waste Business Journal
WERF	Water Environment Research Federation
WFF	World Fab Forecast (previously WFW, World Fab Watch)
WGC	World Gas Conference
WIP	Waste in place
WMO	World Meteorological Organization
WMS	Waste management systems
WTE	Waste-to-energy
WW	Wastewater
WWTP	Wastewater treatment plant
ZEVs	Zero emissions vehicles

6.7. Chemical Formulas

Table A-283: Guide to Chemical Formulas

Table A-203: Guide Ly Cilei	nicai rurniulas
Symbol	Name
Al	Aluminum
Al ₂ O ₃	Aluminum Oxide
Br	Bromine
C	Carbon
CH₄	Methane
C ₂ H ₆	Ethane
C ₃ H ₈	Propane
CF ₄	Perfluoromethane
C ₂ F ₆	Perfluoroethane, hexafluoroethane
c-C ₃ F ₆	Perfluorocyclopropane
C ₃ F ₈	Perfluoropropane
c-C4F8	Perfluorocyclobutane
C ₄ F ₁₀	Perfluorobutane
C ₅ F ₁₂	Perfluoropentane
C ₆ F ₁₄	Perfluorohexane
CF₃I	Trifluoroiodomethane
CFCl₃	Trichlorofluoromethane (CFC-11)
CF ₂ Cl ₂	Dichlorodifluoromethane (CFC-12)
CF ₃ Cl	Chlorotrifluoromethane (CFC-13)
C ₂ F ₃ Cl ₃	Trichlorotrifluoroethane (CFC-113)*
CCI3CF3	CFC-113a*
C ₂ F ₄ Cl ₂	Dichlorotetrafluoroethane (CFC-114)
C ₂ F ₅ Cl	Chloropentafluoroethane (CFC-115)
CHCl₂F	HCFC-21
CHF ₂ Cl	Chlorodifluoromethane (HCFC-22)
C ₂ F ₃ HCl ₂	HCFC-123
C ₂ F ₄ HCI	HCFC-124
C ₂ FH ₃ Cl ₂	HCFC-141b
C ₂ H ₃ F ₂ Cl	HCFC-142b
CF ₃ CF ₂ CHCl ₂	HCFC-225ca
CCIF ₂ CF ₂ CHCIF	HCFC-225cb
CCl ₄	Carbon tetrachloride
CHCICCI ₂	Trichloroethylene
CCI ₂ CCI ₂	Perchloroethylene, tetrachloroethene
CH₃CI	Methylchloride
CH ₃ CCI ₃	Methylchloroform
CH ₂ Cl ₂	Methylenechloride
CHCl₃	Chloroform, trichloromethane
CHF₃	HFC-23
CH ₂ F ₂	HFC-32
CH₃F	HFC-41
C ₂ HF ₅	HFC-125
$C_2H_2F_4$	HFC-134
CH ₂ FCF ₃	HFC-134a
$C_2H_3F_3$	HFC-143*
$C_2H_3F_3$	HFC-143a*
CH ₂ FCH ₂ F	HFC-152*
C ₂ H ₄ F ₂	HFC-152a*
CH ₃ CH ₂ F	HFC-161
C ₃ HF ₇	HFC-227ea
CF ₃ CF ₂ CH ₂ F	HFC-236cb
CF ₃ CHFCHF ₂	HFC-236ea
C3H2F6	HFC-236fa
$C_3H_3F_5$	HFC-245ca

CHF ₂ CH ₂ CF ₃	HFC-245fa
CF ₃ CH ₂ CF ₂ CH ₃	HFC-365mfc
C ₅ H ₂ F ₁₀	HFC-43-10mee
CF ₃ OCHF ₂	HFE-125
CF ₂ HOCF ₂ H	HFE-134
CH ₃ OCF ₃	HFE-143a
CF ₃ CHFOCF ₃	HFF-227ea
CF ₃ CHClOCHF ₂	HCEE-235da2
CF ₃ CHEOCHE ₂	HEF-236ea2
CF ₃ CH ₂ OCF ₃	HEE-236fa
CF ₃ CF ₂ OCH ₃	HEE-245cb2
CHF2CH2OCF3	HFF-245fa1
CE ₃ CH ₂ OCHE ₂	HFF-245fa2
	HEE-254ch2
CE ₂ CH ₂ OCH ₂	HEE-263fb2
	HEE_329mcc2
	HEE-338mcf2
	HEE-347mcc3
	HEE_347mcf2
	HEE-356mac3
	HEE 3560003
	HEE 256nof2
	HEE 256 port2
	HEE 265mof2
	HEE 274 moto
	ПГЕ-3/4рсі2 ЦГГ 7100
	HFO-1234Ze(E)
	HCFO-1233Zd(E)
	H-Galden 1040x
	HG-10
	HG-01 Direction di ette en
	Dimetnyi etner
	Dibromometnane
	Dibromocniorometnane
	Bromodifluoromethane
CH ₃ Br	Methylbromide
CF2BrCl	Bromodichloromethane (Halon 1211)
CF3Br(CBrF3)	Bromotrifluoromethane (Halon 1301)
CF3I	FIC-1311
CO	Carbon monoxide
	Carbon dioxide
CaCO ₃	Calcium carbonate, Limestone
CaMg(CO ₃) ₂	Dolomite
CaO	Calcium oxide, Lime
CI	atomic Chlorine
F	Fluorine
Fe	Iron
Fe ₂ O ₃	Ferric Oxide
FeSi	Ferrosilicon
GaAs	Gallium Arsenide
H, H2	atomic Hydrogen, molecular Hydrogen
H2U	vvater
H2U2	Hydrogen peroxide
UH	
IN, IN2	atomic Nitrogen, molecular Nitrogen

NH ₃	Ammonia
NH4 ⁺	Ammonium ion
HNO ₃	Nitric acid
MgO	Magnesium oxide
NF3	Nitrogen trifluoride
N ₂ O	Nitrous oxide
NO	Nitric oxide
NO ₂	Nitrogen dioxide
NO ₃	Nitrate radical
NOx	Nitrogen oxides
Na	Sodium
Na ₂ CO ₃	Sodium carbonate, soda ash
Na ₃ AIF ₆	Synthetic cryolite
O, O ₂	atomic Oxygen, molecular Oxygen
O ₃	Ozone
S	atomic Sulfur
H ₂ SO ₄	Sulfuric acid
SF ₆	Sulfur hexafluoride
SF5CF3	Trifluoromethylsulphur pentafluoride
SO ₂	Sulfur dioxide
Si	Silicon
SiC	Silicon carbide
SiO ₂	Quartz

* Distinct isomers.

References

- EIA (2018) *Monthly Energy Review, February 2018.* Energy Information Administration, U.S. Department of Energy, Washington, DC. DOE/EIA-0035(2018/2). February 2018.
- EIA (2007) *Emissions of Greenhouse Gases in the United States 2006, Draft Report.* Office of Integrated Analysis and Forecasting, Energy Information Administration, U.S. Department of Energy, Washington, DC. DOE-EIA-0573(2006).
- EIA (1993) *State Energy Data Report 1992*, DOE/EIA-0214(93), Energy Information Administration, U.S. Department of Energy. Washington, DC. December.
- EPA (2016) "1970-2016 Average annual emissions, all criteria pollutants in MS Excel." National Emissions Inventory (NEI) Air Pollutant Emissions Trends Data. Office of Air Quality Planning and Standards. Last Modified December 2016. Available online at: https://www.epa.gov/air-emissions-inventories/air-pollutant-emissions-trends-data.
- EPA (2003) E-mail correspondence. Air pollutant data. Office of Air Pollution to the Office of Air Quality Planning and Standards, U.S. Environmental Protection Agency (EPA). December 22, 2003.
- IPCC (2013) Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change. [Stocker, T.F., D. Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex and P.M. Midgley (eds.)]. Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 1535 pp.
- IPCC (2007) Climate Change 2007: The Physical Science Basis. Contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change. S. Solomon, D. Qin, M. Manning, Z. Chen, M. Marquis, K.B. Averyt, M. Tignor and H.L. Miller (eds.). Cambridge University Press. Cambridge, United Kingdom 996 pp.
- IPCC (1996) Climate Change 1995: The Science of Climate Change. Intergovernmental Panel on Climate Change, J.T. Houghton, L.G. Meira Filho, B.A. Callander, N. Harris, A. Kattenberg, and K. Maskell (eds.). Cambridge University Press. Cambridge, United Kingdom.