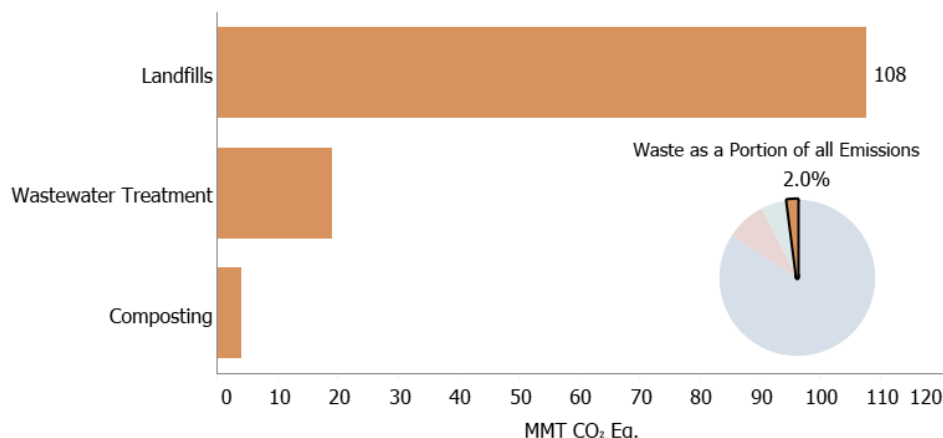


# 7. Waste

Waste management and treatment activities are sources of greenhouse gas emissions (see Figure 7-1). Landfills accounted for approximately 16.2 percent of total U.S. anthropogenic methane (CH<sub>4</sub>) emissions in 2017, the third largest contribution of any CH<sub>4</sub> source in the United States. Additionally, wastewater treatment and composting of organic waste accounted for approximately 2.2 percent and 0.3 percent of U.S. CH<sub>4</sub> emissions, respectively. Nitrous oxide (N<sub>2</sub>O) emissions from the discharge of wastewater treatment effluents into aquatic environments were estimated, as were N<sub>2</sub>O emissions from the treatment process itself. Nitrous oxide emissions from composting were also estimated. Together, these waste activities account for 1.9 percent of total U.S. N<sub>2</sub>O emissions. Nitrogen oxides (NO<sub>x</sub>), carbon monoxide (CO), and non-CH<sub>4</sub> volatile organic compounds (NMVOCs) are emitted by waste activities, and are addressed separately at the end of this chapter. A summary of greenhouse gas emissions from the Waste chapter is presented in Table 7-1 and Table 7-2.

**Figure 7-1: 2017 Waste Chapter Greenhouse Gas Sources (MMT CO<sub>2</sub> Eq.)**



Overall, in 2017, waste activities generated emissions of 131.0 MMT CO<sub>2</sub> Eq., or 2.0 percent of total U.S. greenhouse gas emissions.<sup>1</sup>

**Table 7-1: Emissions from Waste (MMT CO<sub>2</sub> Eq.)**

Gas/Source	1990	2005	2013	2014	2015	2016	2017
<b>CH<sub>4</sub></b>	<b>195.2</b>	<b>148.7</b>	<b>129.3</b>	<b>129.0</b>	<b>127.9</b>	<b>124.4</b>	<b>124.2</b>
Landfills	179.6	131.4	112.9	112.5	111.2	108.0	107.7

<sup>1</sup> Emissions reported in the Waste chapter for landfills and wastewater treatment include those from all 50 states, including Hawaii and Alaska, as well as from U.S. Territories to the extent those waste management activities are occurring. Emissions for composting include all 50 states, including Hawaii and Alaska, but not U.S. Territories. Composting emissions from U.S. Territories are assumed to be small.

Wastewater Treatment	15.3	15.5	14.4	14.4	14.6	14.3	14.3
Composting	0.4	1.9	2.0	2.1	2.1	2.1	2.2
<b>N<sub>2</sub>O</b>	<b>3.7</b>	<b>6.1</b>	<b>6.5</b>	<b>6.6</b>	<b>6.7</b>	<b>6.8</b>	<b>6.9</b>
Wastewater Treatment	3.4	4.4	4.7	4.8	4.8	4.9	5.0
Composting	0.3	1.7	1.8	1.9	1.9	1.9	1.9
<b>Total</b>	<b>199.0</b>	<b>154.8</b>	<b>135.8</b>	<b>135.6</b>	<b>134.5</b>	<b>131.2</b>	<b>131.0</b>

Note: Totals may not sum due to independent rounding.

1 **Table 7-2: Emissions from Waste (kt)**

Gas/Source	1990	2005	2013	2014	2015	2016	2017
<b>CH<sub>4</sub></b>	<b>7,809</b>	<b>5,949</b>	<b>5,173</b>	<b>5,160</b>	<b>5,115</b>	<b>4,975</b>	<b>4,966</b>
Landfills	7,182	5,256	4,517	4,502	4,448	4,319	4,309
Wastewater Treatment	612	618	574	575	582	571	571
Composting	15	75	81	84	85	85	86
<b>N<sub>2</sub>O</b>	<b>12</b>	<b>20</b>	<b>22</b>	<b>22</b>	<b>22</b>	<b>23</b>	<b>23</b>
Wastewater Treatment	11	15	16	16	16	16	17
Composting	1	6	6	6	6	6	6

Note: Totals may not sum due to independent rounding.

2 Carbon dioxide (CO<sub>2</sub>), CH<sub>4</sub>, and N<sub>2</sub>O emissions from the incineration of waste are accounted for in the Energy  
3 sector rather than in the Waste sector because almost all incineration of municipal solid waste (MSW) in the United  
4 States occurs at waste-to-energy facilities where useful energy is recovered. Similarly, the Energy sector also  
5 includes an estimate of emissions from burning waste tires and hazardous industrial waste, because virtually all of  
6 the combustion occurs in industrial and utility boilers that recover energy. The incineration of waste in the United  
7 States in 2017 resulted in 11.1 MMT CO<sub>2</sub> Eq. emissions, more than half of which is attributable to the combustion  
8 of plastics. For more details on emissions from the incineration of waste, see Section 7.4.

#### 9 **Box 7-1: Methodological Approach for Estimating and Reporting U.S. Emissions and Removals**

10 In following the United Nations Framework Convention on Climate Change (UNFCCC) requirement under Article  
11 4.1 to develop and submit national greenhouse gas emission inventories, the emissions and removals presented in  
12 this report and this chapter, are organized by source and sink categories and calculated using internationally-  
13 accepted methods provided by the Intergovernmental Panel on Climate Change (IPCC) in the *2006 IPCC Guidelines*  
14 *for National Greenhouse Gas Inventories (2006 IPCC Guidelines)*. Additionally, the calculated emissions and  
15 removals in a given year for the United States are presented in a common manner in line with the UNFCCC  
16 reporting guidelines for the reporting of inventories under this international agreement. The use of consistent  
17 methods to calculate emissions and removals by all nations providing their inventories to the UNFCCC ensures that  
18 these reports are comparable. The presentation of emissions and sinks provided in this Inventory do not preclude  
19 alternative examinations, but rather, this Inventory presents emissions and removals in a common format consistent  
20 with how countries are to report Inventories under the UNFCCC. The report itself, and this chapter, follows this  
21 standardized format, and provides an explanation of the application of methods used to calculate emissions and  
22 removals.

#### 24 **Box 7-2: Waste Data from EPA's Greenhouse Gas Reporting Program**

On October 30, 2009, the U.S. Environmental Protection Agency (EPA) published a rule requiring annual reporting of greenhouse gas data from large greenhouse gas emission sources in the United States. Implementation of the rule, codified at 40 CFR Part 98, is referred to as EPA's Greenhouse Gas Reporting Program (GHGRP). The rule applies to direct greenhouse gas emitters, fossil fuel suppliers, industrial gas suppliers, and facilities that inject CO<sub>2</sub> underground for sequestration or other reasons and requires reporting by sources or suppliers in 41 industrial categories. Annual reporting is at the facility level, except for certain suppliers of fossil fuels and industrial greenhouse gases. Data reporting by affected facilities includes the

reporting of emissions from fuel combustion at that affected facility. In general, the threshold for reporting is 25,000 metric tons or more of CO<sub>2</sub> Eq. per year.

EPA presents the data collected by its GHGRP through a data publication tool that allows data to be viewed in several formats including maps, tables, charts and graphs for individual facilities or groups of facilities.<sup>2</sup>

EPA's GHGRP dataset and the data presented in this Inventory are complementary. The GHGRP dataset continues to be an important resource for the Inventory, providing not only annual emissions information, but also other annual information, such as activity data and emission factors that can improve and refine national emission estimates and trends over time. GHGRP data also allow EPA to disaggregate national inventory estimates in new ways that can highlight differences across regions and sub-categories of emissions, along with enhancing application of QA/QC procedures and assessment of uncertainties.

EPA uses annual GHGRP data in a number of categories to improve the national estimates presented in this Inventory consistent with IPCC guidelines. Within the Waste Chapter, EPA uses directly reported GHGRP data for net CH<sub>4</sub> emissions from MSW landfills for the years 2010 to 2017 of the Inventory. This data is also used to back-cast emissions from MSW landfills for the years 2005 to 2009.

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## 7.1 Landfills (CRF Source Category 5A1)

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In the United States, solid waste is managed by landfilling, recovery through recycling or composting, and combustion through waste-to-energy facilities. Disposing of solid waste in modern, managed landfills is the most commonly used waste management technique in the United States. More information on how solid waste data are collected and managed in the United States is provided in Box 7-3. The municipal solid waste (MSW) and industrial waste landfills referred to in this section are all modern landfills that must comply with a variety of regulations as discussed in Box 7-3. Disposing of waste in illegal dumping sites is not considered to have occurred in years later than 1980 and these sites are not considered to contribute to net emissions in this section for the timeframe of 1990 to the current Inventory year. MSW landfills, or sanitary landfills, are sites where MSW is managed to prevent or minimize health, safety, and environmental impacts. Waste is deposited in different cells and covered daily with soil; many have environmental monitoring systems to track performance, collect leachate, and collect landfill gas. Industrial waste landfills are constructed in a similar way as MSW landfills, but are used to dispose of industrial solid waste, such as RCRA Subtitle D wastes (e.g., non-hazardous industrial solid waste defined in Title 40 of the Code of Federal Regulations or CFR in section 257.2), commercial solid wastes, or conditionally exempt small-quantity generator wastes (EPA 2016a).

After being placed in a landfill, organic waste (such as paper, food scraps, and yard trimmings) is initially decomposed by aerobic bacteria. After the oxygen has been depleted, the remaining waste is available for consumption by anaerobic bacteria, which break down organic matter into substances such as cellulose, amino acids, and sugars. These substances are further broken down through fermentation into gases and short-chain organic compounds that form the substrates for the growth of methanogenic bacteria. These methane (CH<sub>4</sub>) producing anaerobic bacteria convert the fermentation products into stabilized organic materials and biogas consisting of approximately 50 percent biogenic carbon dioxide (CO<sub>2</sub>) and 50 percent CH<sub>4</sub>, by volume. Landfill biogas also contains trace amounts of non-methane organic compounds (NMOC) and volatile organic compounds (VOC) that either result from decomposition byproducts or volatilization of biodegradable wastes (EPA 2008).

Methane and CO<sub>2</sub> are the primary constituents of landfill gas generation and emissions. However, the *2006 IPCC Guidelines* set an international convention to not report biogenic CO<sub>2</sub> from activities in the Waste sector (IPCC 2006). Net carbon dioxide flux from carbon stock changes in landfills are estimated and reported under the Land Use, Land-Use Change, and Forestry (LULUCF) sector (see Chapter 6 of this Inventory). Additionally, emissions of NMOC and VOC are not estimated because they are emitted in trace amounts. Nitrous oxide (N<sub>2</sub>O) emissions from

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<sup>2</sup> See <[http://www.ipcc-nggip.iges.or.jp/public/tb/TFI\\_Technical\\_Bulletin\\_1.pdf](http://www.ipcc-nggip.iges.or.jp/public/tb/TFI_Technical_Bulletin_1.pdf)>.

1 the disposal and application of sewage sludge on landfills are also not explicitly modeled as part of greenhouse gas  
2 emissions from landfills. Nitrous oxide emissions from sewage sludge applied to landfills as a daily cover or for  
3 disposal are expected to be relatively small because the microbial environment in an anaerobic landfill is not very  
4 conducive to the nitrification and denitrification processes that result in N<sub>2</sub>O emissions. Furthermore, the *2006 IPCC*  
5 *Guidelines* did not include a methodology for estimating N<sub>2</sub>O emissions from solid waste disposal sites “because  
6 they are not significant.” Therefore, only CH<sub>4</sub> generation and emissions are estimated for landfills under the Waste  
7 sector.

8 Methane generation and emissions from landfills are a function of several factors, including: (1) the total amount  
9 and composition of waste-in-place, which is the total waste landfilled annually over the operational lifetime of a  
10 landfill; (2) the characteristics of the landfill receiving waste (e.g., size, climate, cover material); (3) the amount of  
11 CH<sub>4</sub> that is recovered and either flared or used for energy purposes; and (4) the amount of CH<sub>4</sub> oxidized as the  
12 landfill gas – that is not collected by a gas collection system – passes through the cover material into the atmosphere.  
13 Each landfill has unique characteristics, but all managed landfills employ similar operating practices, including the  
14 application of a daily and intermediate cover material over the waste being disposed of in the landfill to prevent odor  
15 and reduce risks to public health. Based on recent literature, the specific type of cover material used can affect the  
16 rate of oxidation of landfill gas (RTI 2011). The most commonly used cover materials are soil, clay, and sand.  
17 Some states also permit the use of green waste, tarps, waste derived materials, sewage sludge or biosolids, and  
18 contaminated soil as a daily cover. Methane production typically begins within the first year after the waste is  
19 disposed of in a landfill and will continue for 10 to 60 years or longer as the degradable waste decomposes over  
20 time.

21 In 2017, landfill CH<sub>4</sub> emissions were approximately 107.7 MMT CO<sub>2</sub> Eq. (4,309 kt), representing the third largest  
22 source of CH<sub>4</sub> emissions in the United States, behind enteric fermentation and natural gas systems. Emissions from  
23 MSW landfills accounted for approximately 95 percent of total landfill emissions, while industrial waste landfills  
24 accounted for the remainder. Estimates of operational MSW landfills in the United States have ranged from 1,700 to  
25 2,000 facilities (EPA 2018a; EPA 2018c; Waste Business Journal [WBJ] 2016; WBJ 2010). More recently, the  
26 Environment Research & Education Foundation (EREF) conducted a nationwide analysis of MSW management and  
27 counted 1,540 operational MSW landfills in 2013 (EREF 2016). Conversely, there are approximately 3,200 MSW  
28 landfills in the United States that have been closed since 1980 (for which a closure data is known, (EPA 2018a; WBJ  
29 2010). While the number of active MSW landfills has decreased significantly over the past 20 years, from  
30 approximately 6,326 in 1990 to as few as 1,540 in the 2013, the average landfill size has increased (EREF 2016;  
31 EPA 2018b; BioCycle 2010). With regard to industrial waste landfills, the WBJ database (WBJ 2016) includes  
32 approximately 1,200 landfills accepting industrial and/or construction and demolition debris for 2016 (WBJ 2016).  
33 Only 172 facilities with industrial waste landfills met the reporting threshold under Subpart TT (Industrial Waste  
34 Landfills) of EPA’s Greenhouse Gas Reporting Program (GHGRP), indicating that there may be several hundred  
35 industrial waste landfills that are not required to report under EPA’s GHGRP.

36 The annual amount of MSW generated and subsequently disposed in MSW landfills varies annually and depends on  
37 several factors (e.g., the economy, consumer patterns, recycling and composting programs, inclusion in a garbage  
38 collection service). The estimated annual quantity of waste placed in MSW landfills increased 10 percent from  
39 approximately 205 MMT in 1990 to 226 MMT in 2000 and then decreased by 8.8 percent to 206 MMT in 2017 (see  
40 Annex 3.14, Table A-253). The total amount of MSW generated is expected to increase as the U.S. population  
41 continues to grow, but the percentage of waste landfilled may decline due to increased recycling and composting  
42 practices. Net CH<sub>4</sub> emissions from MSW landfills have decreased since 1990 (see Table 7-3 and Table 7-4).

43 The estimated quantity of waste placed in industrial waste landfills (from the pulp and paper, and food processing  
44 sectors) has remained relatively steady since 1990, ranging from 9.7 MMT in 1990 to 10.2 MMT in 2017 (see  
45 Annex 3.14, Table A-253). CH<sub>4</sub> emissions from industrial waste landfills have also remained at similar levels  
46 recently, ranging from 14.3 MMT in 2005 to 15.9 MMT in 2017 when accounting for both CH<sub>4</sub> generation and  
47 oxidation.

48 EPA’s Landfill Methane Outreach Program (LMOP) collects information on landfill gas energy projects currently  
49 operational or under construction throughout the United States. LMOP’s project and technical database contains  
50 certain information on the gas collection and control systems in place at landfills that are a part of the program,  
51 which can include the amount of landfill gas collected and flared. In 2017, LMOP identified 15 new landfill gas-to-  
52 energy (LFGE) projects (EPA 2018a) that began operation. While the amount of landfill gas collected and

1 combusted continues to increase, the rate of increase in collection and combustion no longer exceeds the rate of  
 2 additional CH<sub>4</sub> generation from the amount of organic MSW landfilled as the U.S. population grows (EPA 2018b).  
 3 Landfill gas collection and control is not accounted for at industrial waste landfills in this chapter (see the  
 4 Methodology discussion for more information).

5 **Table 7-3: CH<sub>4</sub> Emissions from Landfills (MMT CO<sub>2</sub> Eq.)**

Activity	1990	2005	2013	2014	2015	2016	2017
MSW CH <sub>4</sub> Generation	205.3	-	-	-	-	-	-
Industrial CH <sub>4</sub> Generation	12.1	15.9	16.5	16.6	16.6	16.6	16.6
MSW CH <sub>4</sub> Recovered	(17.9)	-	-	-	-	-	-
MSW CH <sub>4</sub> Oxidized	(18.7)	-	-	-	-	-	-
Industrial CH <sub>4</sub> Oxidized	(1.2)	(1.6)	(1.7)	(1.7)	(1.7)	(1.7)	(1.7)
MSW net CH <sub>4</sub> Emissions (GHGRP)	-	117.1	98.1	97.6	96.3	93.0	92.8
<b>Total</b>	<b>179.6</b>	<b>131.4</b>	<b>112.9</b>	<b>112.5</b>	<b>111.2</b>	<b>108.0</b>	<b>107.7</b>

“-” Not applicable due to methodology change.

Note: Totals may not sum due to independent rounding. Parentheses indicate negative values. For years 1990 to 2004, the Inventory methodology uses the first order decay methodology. A methodological change occurs in year 2005. For years 2005 to 2017, directly reported net CH<sub>4</sub> emissions from the GHGRP data plus a scale-up factor are used to account for emissions from landfill facilities that are not subject to the GHGRP. These data incorporate CH<sub>4</sub> recovered and oxidized. As such, CH<sub>4</sub> generation, CH<sub>4</sub> recovery, and CH<sub>4</sub> oxidized are not calculated separately for 2005 to 2017. See the Time-Series Consistency section of this chapter for more information.

6 **Table 7-4: CH<sub>4</sub> Emissions from Landfills (kt)**

Activity	1990	2005	2013	2014	2015	2016	2017
MSW CH <sub>4</sub> Generation	8,214	-	-	-	-	-	-
Industrial CH <sub>4</sub> Generation	484	636	661	662	663	664	665
MSW CH <sub>4</sub> Recovered	(718)	-	-	-	-	-	-
MSW CH <sub>4</sub> Oxidized	(750)	-	-	-	-	-	-
Industrial CH <sub>4</sub> Oxidized	(48)	(64)	(66)	(66)	(66)	(66)	(67)
MSW net CH <sub>4</sub> Emissions (GHGRP)	-	4,684	3,923	3,906	3,851	3,722	3,711
<b>Total</b>	<b>7,182</b>	<b>5256</b>	<b>4,517</b>	<b>4,502</b>	<b>4,448</b>	<b>4,319</b>	<b>4,309</b>

“-” Not applicable due to methodology change.

Note: Totals may not sum due to independent rounding. Parentheses indicate negative values. For years 1990 to 2004, the Inventory methodology uses the first order decay methodology. A methodological change occurs in year 2005. For years 2005 to 2017, directly reported net CH<sub>4</sub> emissions from the GHGRP data plus a scale-up factor are used to account for emissions from landfill facilities that are not subject to the GHGRP. These data incorporate CH<sub>4</sub> recovered and oxidized. As such, CH<sub>4</sub> generation and CH<sub>4</sub> recovery are not calculated separately. See the Time-Series Consistency section of this chapter for more information.

## 7 Methodology

### 8 Methodology Applied for MSW Landfills

9 Methane emissions from landfills can be estimated using two primary methods. The first method uses the first order  
 10 decay (FOD) model as described by the 2006 IPCC Guidelines to estimate CH<sub>4</sub> generation. The amount of CH<sub>4</sub>  
 11 recovered and combusted from MSW landfills is subtracted from the CH<sub>4</sub> generation and is then adjusted with an  
 12 oxidation factor. The oxidation factor represents the amount of CH<sub>4</sub> in a landfill that is oxidized to CO<sub>2</sub> as it passes  
 13 through the landfill cover (e.g., soil, clay, geomembrane). This method is presented below and is similar to Equation  
 14 HH-5 in 40 CFR Part 98.343 for MSW landfills, and Equation TT-6 in 40 CFR Part 98.463 for industrial waste  
 15 landfills.

$$\text{CH}_{4,\text{Solid Waste}} = [\text{CH}_{4,\text{MSW}} + \text{CH}_{4,\text{Ind}} - \text{R}] - \text{Ox}$$

17 where,

- 1 CH<sub>4,Solid Waste</sub> = Net CH<sub>4</sub> emissions from solid waste
- 2 CH<sub>4,MSW</sub> = CH<sub>4</sub> generation from MSW landfills
- 3 CH<sub>4,Ind</sub> = CH<sub>4</sub> generation from industrial waste landfills
- 4 R = CH<sub>4</sub> recovered and combusted (only for MSW landfills)
- 5 OX = CH<sub>4</sub> oxidized from MSW and industrial waste landfills before release to the atmosphere

6 The second method used to calculate CH<sub>4</sub> emissions from landfills, also called the back-calculation method, is based  
 7 on directly measured amounts of recovered CH<sub>4</sub> from the landfill gas and is expressed below and by Equation HH-8  
 8 in 40 CFR Part 98.343. The two parts of the equation consider the portion of CH<sub>4</sub> in the landfill gas that is not  
 9 collected by the landfill gas collection system, and the portion that is collected. First, the recovered CH<sub>4</sub> is adjusted  
 10 with the collection efficiency of the gas collection and control system and the fraction of hours the recovery system  
 11 operated in the calendar year. This quantity represents the amount of CH<sub>4</sub> in the landfill gas that is not captured by  
 12 the collection system; this amount is then adjusted for oxidation. The second portion of the equation adjusts the  
 13 portion of CH<sub>4</sub> in the collected landfill gas with the efficiency of the destruction device(s), and the fraction of hours  
 14 the destruction device(s) operated during the year.

$$15 \quad CH_{4,Solid\ Waste} = \left[ \left( \frac{R}{CE \times f_{REC}} - R \right) \times (1 - OX) + R \times (1 - (DE \times f_{Dest})) \right]$$

16 where,

- 17 CH<sub>4,Solid Waste</sub> = Net CH<sub>4</sub> emissions from solid waste
- 18
- 19 R = Quantity of recovered CH<sub>4</sub> from Equation HH-4 of EPA’s GHGRP
- 20 CE = Collection efficiency estimated at the landfill, considering system coverage, operation,  
 21 and cover system materials from Table HH-3 of EPA’s GHGRP. If area by soil cover type  
 22 information is not available, the default value of 0.75 should be used. (percent)
- 23 f<sub>REC</sub> = fraction of hours the recovery system was operating (percent)
- 24 OX = oxidation factor (percent)
- 25 DE = destruction efficiency (percent)
- 26 f<sub>Dest</sub> = fraction of hours the destruction device was operating (fraction)
- 27

28 The current Inventory uses both methods to estimate CH<sub>4</sub> emissions across the time series. Prior to the 1990 through  
 29 2015 Inventory, only the FOD method was used. Methodological changes were made to the 1990 through 2015  
 30 Inventory to incorporate higher tier data (i.e., directly reported CH<sub>4</sub> emissions to EPA’s GHGRP), which cannot be  
 31 directly applied to earlier years in the time series without significant bias. The technique used to merge the directly  
 32 reported GHGRP data with the previous methodology is described as the overlap technique in the Time-Series  
 33 Consistency chapter of the *2006 IPCC Guidelines*. Additional details on the technique used is included in the Time  
 34 Series Consistency section of this chapter and a technical memorandum (RTI 2017).

35 A summary of the methodology used to generate the current 1990 through 2017 Inventory estimates for MSW  
 36 landfills is as follows and also illustrated in Annex Figure A-18:

- 37 • **1940 through 1989:** These years are included for historical waste disposal amounts. Estimates of the  
 38 annual quantity of waste landfilled for 1960 through 1988 were obtained from EPA’s *Anthropogenic*  
 39 *Methane Emissions in the United States, Estimates for 1990: Report to Congress* (EPA 1993) and an  
 40 extensive landfill survey by the EPA’s Office of Solid Waste in 1986 (EPA 1988). Although waste placed  
 41 in landfills in the 1940s and 1950s contributes very little to current CH<sub>4</sub> generation, estimates for those  
 42 years were included in the FOD model for completeness in accounting for CH<sub>4</sub> generation rates and are  
 43 based on the population in those years and the per capita rate for land disposal for the 1960s. For the  
 44 Inventory calculations, wastes landfilled prior to 1980 were broken into two groups: wastes disposed in  
 45 managed, anaerobic landfills (Methane Conversion Factor, MCF, of 1) and those disposed in uncategorized  
 46 solid waste disposal waste sites (MCF of 0.6) (IPCC 2006). Uncategorized sites represent those sites for  
 47 which limited information is known about the management practices. All calculations after 1980 assume  
 48 waste is disposed in managed, anaerobic landfills. The FOD method was applied to estimate annual CH<sub>4</sub>  
 49 generation. Methane recovery amounts were then subtracted and the result was then adjusted with a 10  
 50 percent oxidation factor to derive the net emissions estimates.

- 1 • **1990 through 2004:** The Inventory time series begins in 1990. The FOD method is exclusively used for  
2 this group of years. The national total of waste generated (based on state-specific landfill waste generation  
3 data) and a national average disposal factor for 1989 through 2004 were obtained from the State of Garbage  
4 (SOG) survey every two years (i.e., 2002, 2004 as published in BioCycle 2006). In-between years were  
5 interpolated based on population growth. For years 1989 to 2000, directly reported total MSW generation  
6 data were used; for other years, the estimated MSW generation (excluding construction and demolition  
7 waste and inerts) were presented in the reports and used in the Inventory. The FOD method was applied to  
8 estimate annual CH<sub>4</sub> generation. Landfill-specific CH<sub>4</sub> recovery amounts were then subtracted from CH<sub>4</sub>  
9 generation and the result was then adjusted with a 10 percent oxidation factor to derive the net emissions  
10 estimates.
- 11 • **2005 through 2009:** Emissions for these years are estimated using net CH<sub>4</sub> emissions that are reported by  
12 landfill facilities under EPA's GHGRP. Because not all landfills in the United States are required to report  
13 to EPA's GHGRP, a 9 percent scale-up factor is applied to the GHGRP emissions for completeness.  
14 Supporting information, including details on the technique used to estimate emissions for 2005 to 2009 and  
15 to ensure time-series consistency by incorporating the directly reported GHGRP emissions is presented in  
16 Annex 3.14 and in RTI 2018a. A single oxidation factor is not applied to the annual CH<sub>4</sub> generated as is  
17 done for 1990 to 2004 because the GHGRP emissions data are used, which already take oxidation into  
18 account. The GHGRP allows facilities to use varying oxidation factors depending on their facility-specific  
19 calculated CH<sub>4</sub> flux rate (i.e., 0, 10, 25, or 35 percent). The average oxidation factor from the GHGRP  
20 facilities is 19.5 percent.
- 21 • **2010 through 2017:** Directly reported net CH<sub>4</sub> emissions to the GHGRP are used with a 9 percent scale-up  
22 factor to account for landfills that are not required to report to the GHGRP. A combination of the FOD  
23 method and the back-calculated CH<sub>4</sub> emissions were used by the facilities reporting to the GHGRP.  
24 Landfills reporting to the GHGRP without gas collection and control apply the FOD method, while most  
25 landfills with landfill gas collection and control apply the back-calculation method. As noted above,  
26 GHGRP facilities use a variety of oxidation factors. The average oxidation factor from the GHGRP  
27 facilities is 19.5 percent.

28 A detailed discussion of the data sources and methodology used to calculate CH<sub>4</sub> generation and recovery is  
29 provided below. Supporting information, including details on the technique used to ensure time-series consistency  
30 by incorporating the directly reported GHGRP emissions is presented in the Time-Series Consistency section of this  
31 chapter and in Annex 3.14.

## 32 **Description of the Methodology for MSW Landfills as Applied for 1990-2004**

### 33 *National MSW Methane Generation and Disposal Estimates*

34 States and local municipalities across the United States do not consistently track and report quantities of MSW  
35 generated or collected for management, nor do they report end-of-life disposal methods to a centralized system.  
36 Therefore, national MSW landfill waste generation and disposal data are obtained from secondary data, specifically  
37 the SOG surveys, published approximately every two years, with the most recent publication date of 2014. The SOG  
38 survey was the only continually updated nationwide survey of waste disposed in landfills in the United States and  
39 was the primary data source with which to estimate nationwide CH<sub>4</sub> generation from MSW landfills. Currently,  
40 EPA's GHGRP waste disposal data and MSW management data published by EREF are available.

41 The SOG surveys collect data from the state agencies and then apply the principles of mass balance where all MSW  
42 generated is equal to the amount of MSW landfilled, combusted in waste-to-energy plants, composted, and/or  
43 recycled (BioCycle 2006; Shin 2014). This approach assumes that all waste management methods are tracked and  
44 reported to state agencies. Survey respondents are asked to provide a breakdown of MSW generated and managed  
45 by landfilling, recycling, composting, and combustion (in waste-to-energy facilities) in actual tonnages as opposed  
46 to reporting a percent generated under each waste disposal option. The data reported through the survey have  
47 typically been adjusted to exclude non-MSW materials (e.g., industrial and agricultural wastes, construction and  
48 demolition debris, automobile scrap, and sludge from wastewater treatment plants) that may be included in survey  
49 responses. While these wastes may be disposed of in MSW landfills, they are not the primary type of waste material

1 disposed and are typically inert. In the most recent survey, state agencies were asked to provide already filtered,  
2 MSW-only data. Where this was not possible, they were asked to provide comments to better understand the data  
3 being reported. All state disposal data are adjusted for imports and exports across state lines where imported waste is  
4 included in a state's total while exported waste is not. Methodological changes have occurred over the time frame  
5 the SOG survey has been published, and this has affected the fluctuating trends observed in the data (RTI 2013).

6 State-specific landfill MSW generation data and a national average disposal factor for 1989 through 2004 were  
7 obtained from the SOG survey every two years (i.e., 2002, 2004 as published in BioCycle 2006). The landfill  
8 inventory calculations start with hard numbers (where available) as presented in the SOG documentation for the  
9 report years 2002 and 2004. In-between year waste generation is interpolated using the prior and next SOG report  
10 data. For example, waste generated in 2003 = (waste generation in 2002 + waste generation in 2004)/2. The  
11 quantities of waste generated across all states are summed and that value is then used as the nationwide quantity of  
12 waste generated in each year of the time series. The SOG survey is voluntary and not all states provide data in each  
13 survey year. To estimate waste generation for states that did not provide data in any given reporting year, one of the  
14 following methods was used (RTI 2013):

- 15 • For years when a state-specific waste generation rate was available from the previous SOG reporting year  
16 submission, the state-specific waste generation rate for that particular state was used.

17 – or –

- 18 • For years where a state-specific waste generation rate was not available from the previous SOG reporting  
19 year submission, the waste amount is generated using the national average waste generation rate. In other  
20 words, Waste Generated = Reporting Year Population × the National Average Waste Generation Rate
  - 21 ○ The National Average Waste Generation Rate is determined by dividing the total reported waste  
22 generated across the reporting states by the total population for reporting states.
  - 23 ○ This waste generation rate may be above or below the waste generation rate for the non-reporting  
24 states and contributes to the overall uncertainty of the annual total waste generation amounts used  
25 in the model.

26 Use of these methods to estimate solid waste generated by states is a key aspect of how the SOG data was  
27 manipulated and why the results differ for total solid waste generated as estimated by SOG and in the Inventory. In  
28 the early years (2002 data in particular), SOG made no attempt to fill gaps for non-survey responses. For the 2004  
29 data, the SOG team used proxy data (mainly from the WBJ) to fill gaps for non-reporting states and survey  
30 responses.

31 Another key aspect of the SOG survey is that it focuses on MSW-only data. The data states collect for solid waste  
32 typically are representative of total solid waste and not the MSW-only fraction. In the early years of the SOG  
33 survey, most states reported total solid waste rather than MSW-only waste. The SOG team, in response, “filtered”  
34 the state-reported data to reflect the MSW-only portion.

35 This data source also contains the waste generation data reported by states to the SOG survey, which fluctuates from  
36 year to year. Although some fluctuation is expected, for some states, the year-to-year fluctuations are quite  
37 significant (>20 percent increase or decrease in some case) (RTI 2013). The SOG survey reports for these years do  
38 not provide additional explanation for these fluctuations and the source data are not available for further assessment.  
39 Although exact reasons for the large fluctuations are difficult to obtain without direct communication with states,  
40 staff from the SOG team that were contacted speculate that significant fluctuations are present because the particular  
41 state could not gather complete information for waste generation (i.e., they are missing part of recycled and  
42 composted waste data) during a given reporting year. In addition, SOG team staff speculated that some states may  
43 have included C&D and industrial wastes in their previous MSW generation submissions, but made efforts to  
44 exclude that (and other non-MSW categories) in more recent reports (RTI 2013).

45 Recently, the EREF published a report, *MSW Management in the United States*, which includes state-specific  
46 landfill MSW generation and disposal data for 2010 and 2013 using a similar methodology as the SOG surveys  
47 (EREF 2016). Because of this similar methodology, EREF data were used to populate data for years where BioCycle  
48 data was not available when possible. State-specific landfill waste generation data for the years in between the SOG  
49 surveys and EREF report (e.g., 2001, 2003, etc.) were either interpolated or extrapolated based on the SOG or EREF  
50 data and the U.S. Census population data (U.S. Census Bureau 2018).



1 Estimates of the quantity of waste landfilled from 1989 to 2004 are determined by applying an average national  
2 waste disposal factor to the total amount of waste generated (i.e., the SOG data). A national average waste disposal  
3 factor is determined for each year an SOG survey is published and equals the ratio of the total amount of waste  
4 landfilled in the United States to the total amount of waste generated in the United States. The waste disposal factor  
5 is interpolated or extrapolated for the years in-between the SOG surveys, as is done for the amount of waste  
6 generated for a given survey year.

7 The 2006 IPCC Guidelines recommend at least 50 years of waste disposal data to estimate CH<sub>4</sub> emissions. Estimates  
8 of the annual quantity of waste landfilled for 1960 through 1988 were obtained from EPA's *Anthropogenic Methane*  
9 *Emissions in the United States, Estimates for 1990: Report to Congress* (EPA 1993) and an extensive landfill survey  
10 by the EPA's Office of Solid Waste in 1986 (EPA 1988). Although waste placed in landfills in the 1940s and 1950s  
11 contributes very little to current CH<sub>4</sub> generation, estimates for those years were included in the FOD model for  
12 completeness in accounting for CH<sub>4</sub> generation rates and are based on the population in those years and the per  
13 capita rate for land disposal for the 1960s. For calculations in the current Inventory, wastes landfilled prior to 1980  
14 were broken into two groups: wastes disposed in landfills (MCF of 1) and those disposed in uncategorized site as  
15 (MCF of 0.6). All calculations after 1980 assume waste is disposed in managed, modern landfills. See Annex 3.14  
16 for more details.

17 In the current Inventory methodology, the MSW generation and disposal data are no longer used to estimate CH<sub>4</sub>  
18 emissions for the years 2005 to 2017 because EPA's GHGRP emissions data are now used for those years.

### 19 *National Landfill Gas Recovery Estimates for MSW Landfills*

20 The estimated landfill gas recovered per year (R) at MSW landfills for 1990 to 2004 was based on a combination of  
21 four databases and includes recovery from flares and/or landfill gas-to-energy (LFGE) projects:

- 22 • EPA's GHGRP dataset for MSW landfills (EPA 2015a);<sup>3</sup>
- 23 • A database developed by the Energy Information Administration (EIA) for the voluntary reporting of  
24 greenhouse gases (EIA 2007);
- 25 • A database of LFGE projects that is primarily based on information compiled by the EPA LMOP (EPA  
26 2016b);<sup>4</sup> and
- 27 • The flare vendor database (contains updated sales data collected from vendors of flaring equipment).

28 The same landfill may be included one or more times across these four databases. To avoid double- or triple-  
29 counting CH<sub>4</sub> recovery, the landfills across each database were compared and duplicates identified. A hierarchy of  
30 recovery data is used based on the certainty of the data in each database. In summary, the GHGRP > EIA > LFGE >  
31 flare vendor database. The rationale for this hierarchy is described below.

32 EPA's GHGRP MSW landfills database was first introduced as a data source for landfill gas recovery in the 1990 to  
33 2013 Inventory. EPA's GHGRP MSW landfills database contains facility-reported data that undergoes rigorous  
34 verification, thus it is considered to contain the least uncertain data of the four CH<sub>4</sub> recovery databases. However, as  
35 mentioned earlier, this database is unique in that it only contains a portion of the landfills in the United States  
36 (although, presumably the highest emitters since only those landfills that meet a certain CH<sub>4</sub> generation threshold  
37 must report) and only contains data for 2010 and later. In the current Inventory methodology, CH<sub>4</sub> recovery for 1990  
38 to 2004 for facilities reporting to EPA's GHGRP has been estimated using the directly reported emissions for those  
39 facilities from 2010 to 2015, and an Excel forecasting function so that the GHGRP data source can be applied to  
40 earlier years in the time series. Prior to 2005, if a landfill in EPA's GHGRP was also in the LFGE or EIA databases,  
41 the landfill gas project information, specifically the project start year, from either the LFGE or EIA databases was  
42 used as the cutoff year for the estimated CH<sub>4</sub> recovery in the GHGRP database. For example, if a landfill reporting  
43 under EPA's GHGRP was also included in the LFGE database under a project that started in 2002 that is still

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<sup>3</sup> The 2015 GHGRP dataset is used to estimate landfill gas recovery from MSW landfills for the years 1990 to 2004 of the Inventory. This database is no longer updated because the methodology has changed such that the directly reported net methane emissions from the GHGRP are used and landfill gas recovery is not separately estimated.

<sup>4</sup> The LFGE database was not updated for the 1990 to 2017 Inventory because the methodology does not use this database for years 2005 and later, thus the LMOP 2016 database is the most recent year reflected in the LFGE database for the Inventory.

1 operational, the CH<sub>4</sub> recovery data in the GHGRP database for that facility was back-calculated to the year 2002  
2 only.

3 If a landfill in the GHGRP MSW landfills database was also in the EIA, LFGE, and/or flare vendor database, the  
4 avoided emissions were only based on EPA's GHGRP MSW landfills database to avoid double or triple counting  
5 the recovery amounts. In other words, the CH<sub>4</sub> recovery from the same landfill was not included in the total recovery  
6 from the EIA, LFGE, or flare vendor databases.

7 If a landfill in the EIA database was also in the LFGE and/or the flare vendor database, the CH<sub>4</sub> recovery was based  
8 on the EIA data because landfill owners or operators directly reported the amount of CH<sub>4</sub> recovered using gas flow  
9 concentration and measurements, and because the reporting accounted for changes over time.

10 If both the flare data and LFGE recovery data were available for any of the remaining landfills (i.e., not in the EIA  
11 or GHGRP databases), then the avoided emissions were based on the LFGE data, which provides reported landfill-  
12 specific data on gas flow for direct use projects and project capacity (i.e., megawatts) for electricity projects. The  
13 LFGE database is based on the most recent EPA LMOP database (published annually). The remaining portion of  
14 avoided emissions is calculated by the flare vendor database, which estimates CH<sub>4</sub> combusted by flares using the  
15 midpoint of a flare's reported capacity. New flare vendor sales data have not been collected since 2015 because  
16 these data are not used for estimates beyond 2005 in the time series. Given that each LFGE project is likely to also  
17 have a flare, double counting reductions from flares and LFGE projects in the LFGE database was avoided by  
18 subtracting emission reductions associated with LFGE projects for which a flare had not been identified from the  
19 emission reductions associated with flares (referred to as the flare correction factor). A further explanation of the  
20 methodology used to estimate the landfill gas recovered can be found in Annex 3.14.

21 A destruction efficiency of 99 percent was applied to CH<sub>4</sub> recovered to estimate CH<sub>4</sub> emissions avoided due to the  
22 combusting of CH<sub>4</sub> in destruction devices (i.e., flares) in the EIA, LFGE, and flare vendor databases. The median  
23 value of the reported destruction efficiencies to the GHGRP was 99 percent for all reporting years (2010 through  
24 2017). For the other three recovery databases, the 99 percent destruction efficiency value selected was based on the  
25 range of efficiencies (86 to greater than 99 percent) recommended for flares in EPA's *AP-42 Compilation of Air  
26 Pollutant Emission Factors*, Draft Section 2.4, Table 2.4-3 (EPA 2008). A typical value of 97.7 percent was  
27 presented for the non-CH<sub>4</sub> components (i.e., VOC and NMOC) in test results (EPA 2008). An arithmetic average of  
28 98.3 percent and a median value of 99 percent are derived from the test results presented in EPA (2008). Thus, a  
29 value of 99 percent for the destruction efficiency of flares has been used in the Inventory methodology. Other data  
30 sources supporting a 99 percent destruction efficiency include those used to establish New Source Performance  
31 Standards (NSPS) for landfills and in recommendations for shutdown flares used by the EPA LMOP.

### 32 *National MSW Landfill Methane Oxidation Estimates*

33 The amount of CH<sub>4</sub> oxidized by the landfill cover at MSW landfills was assumed to be 10 percent of the CH<sub>4</sub>  
34 generated that is not recovered (IPCC 2006; Mancinelli and McKay 1985; Czepiel et al. 1996) for the years 1990 to  
35 2004.

### 36 *National MSW Net Emissions Estimates*

37 Net CH<sub>4</sub> emissions are calculated by subtracting the CH<sub>4</sub> recovered and CH<sub>4</sub> oxidized from CH<sub>4</sub> generated at MSW  
38 landfills.

## 39 **Description of the Methodology for MSW Landfills as Applied for 2005 to 2009**

40 The Inventory methodology uses directly reported net CH<sub>4</sub> emissions for the 2010 to 2017 reporting years from  
41 EPA's GHGRP to back-cast emissions for 2005 to 2009. The emissions for 2005 to 2009 are recalculated each year  
42 the Inventory is published to account for the additional year of reported data and any revisions that facilities make to  
43 past GHGRP reports. When EPA verifies the greenhouse gas reports, comparisons are made with data submitted in  
44 earlier reporting years and errors may be identified in these earlier year reports. Facility representatives may submit  
45 revised reports for any reporting year in order to correct these errors. Facilities reporting to EPA's GHGRP that do  
46 not have landfill gas collection and control systems use the FOD method. Facilities with landfill gas collection and  
47 control must use both the FOD method and a back-calculation approach. The back-calculation approach starts with

1 the amount of CH<sub>4</sub> recovered and works back through the system to account for gas not collected by the landfill gas  
2 collection and control system (i.e., the collection efficiency).

3 A scale-up factor to account for emissions from landfills that do not report to EPA's GHGRP is also applied  
4 annually. In theory, national MSW landfill emissions should equal the net CH<sub>4</sub> emissions reported to the GHGRP  
5 plus net CH<sub>4</sub> emissions from landfills that do not report to the GHGRP. EPA estimated a scale-up factor of 9  
6 percent. The rationale behind the 9 percent scale-up factor is included in Annex 3.14 and in RTI 2018a.

7 The GHGRP data allows facilities to apply a range of oxidation factors (0.0, 0.10, 0.25, or 0.35) based on the  
8 calculated CH<sub>4</sub> flux at the landfill. Therefore, one oxidation factor is not applied for this time frame, as is done for  
9 1990 to 2004. The average oxidation factor across the GHGRP data is 19.5 percent.

## 10 **Description of the Methodology for MSW Landfills as Applied for 2010 to 2017**

11 Directly reported CH<sub>4</sub> emissions to the GHGRP are used for 2010 to 2017 plus the 9 percent scale-up factor to  
12 account for emissions from landfills that do not report to the GHGRP. The average oxidation factor across the  
13 GHGRP data is 19.5 percent.

## 14 **Description of the First Order Decay Methodology for Industrial Waste Landfills**

15 Emissions from industrial waste landfills are estimated from industrial production data (ERG 2018), waste disposal  
16 factors, and the FOD method. There are currently no data sources that track and report the amount and type of waste  
17 disposed of in the universe of industrial waste landfills in the United States. EPA's GHGRP provides some insight  
18 into waste disposal in industrial waste landfills, but is not comprehensive. Data reported to the GHGRP on industrial  
19 waste landfills suggests that most of the organic waste which would result in methane emissions is disposed at pulp  
20 and paper and food processing facilities. Of the 172 facilities that reported to subpart TT of the GHGRP in 2017, 93  
21 (54 percent) are in the North American Industrial Classification System (NAICS) for Pulp, Paper, and Wood  
22 Products (NAICS 321 and 322) and 12 (7 percent) are in Food Manufacturing (NAICS 311). Based on this limited  
23 information, the Inventory methodology assumes most of the organic waste placed in industrial waste landfills  
24 originates from the food processing (meat, vegetables, fruits) and pulp and paper sectors, thus estimates of industrial  
25 landfill emissions focused on these two sectors. To validate this assumption, the EPA recently conducted an analysis  
26 of data reported to subpart TT of the GHGRP in the 2016 reporting year. Waste streams of facilities reporting to  
27 subpart TT were designated as either relating to food and beverage, pulp and paper, or other based on their primary  
28 NAICS code. The total waste disposed by facilities under each primary NAICS reported in 2016 were calculated in  
29 order to determine that 93 percent of the total organic waste quantity reported under subpart TT is originating from  
30 either the pulp and paper or food and beverage sector (RTI 2018b).

31 The composition of waste disposed of in industrial waste landfills is expected to be more consistent in terms of  
32 composition and quantity than that disposed of in MSW landfills. The amount of waste landfilled is assumed to be a  
33 fraction of production that is held constant over the time series as explained in Annex 3.14.

34 Landfill CH<sub>4</sub> recovery is not accounted for in industrial waste landfills. Data collected through EPA's GHGRP for  
35 industrial waste landfills (Subpart TT) show that only two of the 172 facilities, or 1 percent of facilities, have active  
36 gas collection systems (EPA 2018b). However, because EPA's GHGRP is not a national database and  
37 comprehensive data regarding gas collection systems have not been published for industrial waste landfills,  
38 assumptions regarding a percentage of landfill gas collection systems, or a total annual amount of landfill gas  
39 collected for the non-reporting industrial waste landfills have not been made for the Inventory methodology.

40 The amount of CH<sub>4</sub> oxidized by the landfill cover at industrial waste landfills was assumed to be 10 percent of the  
41 CH<sub>4</sub> generated (IPCC 2006; Mancinelli and McKay 1985; Czepiel et al. 1996) for all years.

## 42 **Uncertainty and Time-Series Consistency**

43 Several types of uncertainty are associated with the estimates of CH<sub>4</sub> emissions from MSW and industrial waste  
44 landfills when the FOD method is applied directly for 1990 to 2004 in the Waste Model and, to some extent, in the  
45 GHGRP methodology. The approach used in the MSW emission estimates assumes that the CH<sub>4</sub> generation  
46 potential (L<sub>0</sub>) and the rate of decay that produces CH<sub>4</sub> from MSW, as determined from several studies of CH<sub>4</sub>  
47 recovery at MSW landfills, are representative of conditions at U.S. MSW landfills. When this top-down approach is

1 applied at the nationwide level, the uncertainties are assumed to be less than when applying this approach to  
2 individual landfills and then aggregating the results to the national level. In other words, the FOD method as applied  
3 in this Inventory is not facility-specific modeling and while this approach may over- or under-estimate CH<sub>4</sub>  
4 generation at some landfills if used at the facility-level, the result is expected to balance out because it is being  
5 applied nationwide.

6 There is a high degree of uncertainty associated with the FOD model, particularly when a homogeneous waste  
7 composition and hypothetical decomposition rates are applied to heterogeneous landfills (IPCC 2006). There is less  
8 uncertainty in EPA's GHGRP data because this methodology is facility-specific, uses directly measured CH<sub>4</sub>  
9 recovery data (when applicable), and allows for a variety of landfill gas collection efficiencies, destruction  
10 efficiencies, and/or oxidation factors to be used.

11 Uncertainty also exists in the scale-up factor applied for years 2005 to 2009 and in the back-casted emissions  
12 estimates for 2005 to 2009. As detailed in RTI (2018a), limited information is available for landfills that do not  
13 report to the GHGRP. RTI developed an initial list of landfills that do not report to the GHGRP with the intent of  
14 quantifying the total waste-in-place for these landfills that would add up to the scale-up factor. Input was provided  
15 by industry, LMOP, and additional EPA support. However, many gaps still exist and assumptions were made for  
16 many landfills in order to estimate the scale-up factor. Additionally, a simple methodology was used to back-cast  
17 emissions for 2005 to 2009 using the GHGRP emissions from 2010 to 2017. This methodology does not factor in  
18 annual landfill to landfill changes in landfill CH<sub>4</sub> generation and recovery. Because of this, an uncertainty factor of  
19 25 percent is applied to emissions for 2005 to 2009.

20 With regard to the time series and as stated in *2006 IPCC Guidelines Volume 1: Chapter 5 Time-Series Consistency*  
21 (IPCC 2006), "the time series is a central component of the greenhouse gas inventory because it provides  
22 information on historical emissions trends and tracks the effects of strategies to reduce emissions at the national  
23 level. All emissions in a time series should be estimated consistently, which means that as far as possible, the time  
24 series should be calculated using the same method and data sources in all years" (IPCC 2006). This chapter  
25 however, recommends against back-casting emissions back to 1990 with a limited set of data and instead provides  
26 guidance on techniques to splice, or join methodologies together. One of those techniques is referred to as the  
27 overlap technique. The overlap technique is recommended when new data becomes available for multiple years.  
28 This was the case with the GHGRP data for MSW landfills, where directly reported CH<sub>4</sub> emissions data became  
29 available for more than 1,200 MSW landfills beginning in 2010. The GHGRP emissions data had to be merged with  
30 emissions from the FOD method to avoid a drastic change in emissions in 2010, when the datasets were combined.  
31 EPA also had to consider that according to IPCC's good practice, efforts should be made to reduce uncertainty in  
32 Inventory calculations and that, when compared to the GHGRP data, the FOD method presents greater uncertainty.

33 In evaluating the best way to combine the two datasets, EPA considered either using the FOD method from 1990 to  
34 2009, or using the FOD method for a portion of that time and back-casting the GHGRP emissions data to a year  
35 where emissions from the two methodologies aligned. Plotting the back-casted GHGRP emissions against the  
36 emissions estimates from the FOD method showed an alignment of the data in 2004 and later years which facilitated  
37 the use of the overlap technique while also reducing uncertainty. Therefore, EPA decided to back-cast the GHGRP  
38 emissions from 2009 to 2005 only, in order to merge the datasets and adhere to the *IPCC Good Practice Guidance*  
39 for ensuring time series consistency.

40 Aside from the uncertainty in estimating landfill CH<sub>4</sub> generation, uncertainty also exists in the estimates of the  
41 landfill gas oxidized at MSW landfills. Facilities directly reporting to EPA's GHGRP can use oxidation factors  
42 ranging from 0 to 35 percent, depending on their facility-specific CH<sub>4</sub> flux. As recommended by the *2006 IPCC*  
43 *Guidelines* for managed landfills, a 10 percent default oxidation factor is applied in the Inventory for both MSW  
44 landfills (those not reporting to the GHGRP and for the years 1990 to 2004 when GHGRP data are not available)  
45 and industrial waste landfills regardless of climate, the type of cover material, and/or presence of a gas collection  
46 system. The number of published field studies measuring the rate of oxidation has increased substantially since the  
47 *2006 IPCC Guidelines* were published and, as discussed in the Potential Improvements section, efforts will continue  
48 to review the literature and revise this value, as appropriate.

49 Another significant source of uncertainty lies with the estimates of CH<sub>4</sub> recovered by flaring and gas-to-energy  
50 projects at MSW landfills that are sourced from the Inventory's CH<sub>4</sub> recovery databases (used for years 1990 to  
51 2004). Four CH<sub>4</sub> recovery databases are used to estimate nationwide CH<sub>4</sub> recovery for MSW landfills for 1990 to  
52 2004; whereas directly reported CH<sub>4</sub> recovery is used for facilities reporting to the GHGRP for years 2005 to 2015.  
53 The GHGRP MSW landfills database was added as a fourth recovery database starting with the 1990 through 2013

1 Inventory report. Relying on multiple databases for a complete picture introduces uncertainty because the coverage  
2 and characteristics of each database differs, which increases the chance of double counting avoided emissions.  
3 Additionally, the methodology and assumptions that go into each database differ. For example, the flare database  
4 assumes the midpoint of each flare capacity at the time it is sold and installed at a landfill; the flare may be  
5 achieving a higher capacity, in which case the flare database would underestimate the amount of CH<sub>4</sub> recovered.

6 The LFGE database was updated annually until 2015. The flare database was populated annually until 2015 by the  
7 voluntary sharing of flare sales data by select vendors, which likely underestimated recovery for landfills not  
8 included in the three other recovery databases used by the Inventory. The EIA database has not been updated since  
9 2006 and has, for the most part, been replaced by the GHGRP MSW landfills database. To avoid double counting  
10 and to use the most relevant estimate of CH<sub>4</sub> recovery for a given landfill, a hierarchical approach is used among the  
11 four databases. GHGRP data and the EIA data are given precedence because facility data were directly reported; the  
12 LFGE data are given second priority because CH<sub>4</sub> recovery is estimated from facility-reported LFGE system  
13 characteristics; and the flare data are given the lowest priority because this database contains minimal information  
14 about the flare, no site-specific operating characteristics, and includes smaller landfills not included in the other  
15 three databases (Bronstein et al. 2012). The coverage provided across the databases most likely represents the  
16 complete universe of landfill CH<sub>4</sub> gas recovery; however, the number of unique landfills between the four databases  
17 does differ.

18 The 2006 IPCC Guidelines default value of 10 percent for uncertainty in recovery estimates was used for two of the  
19 four recovery databases in the uncertainty analysis where metering of landfill gas was in place (for about 64 percent  
20 of the CH<sub>4</sub> estimated to be recovered). This 10 percent uncertainty factor applies to the LFGE database; 12 percent  
21 to the EIA database; and 1 percent for the GHGRP MSW landfills dataset because of the supporting information  
22 provided and rigorous verification process. For flaring without metered recovery data (the flare database), a much  
23 higher uncertainty value of 50 percent is used. The compounding uncertainties associated with the four databases in  
24 addition to the uncertainties associated with the FOD method and annual waste disposal quantities leads to the large  
25 upper and lower bounds for MSW landfills presented in Table 7-5.

26 The lack of landfill-specific information regarding the number and type of industrial waste landfills in the United  
27 States is a primary source of uncertainty with respect to the industrial waste generation and emission estimates. The  
28 approach used here assumes that most of the organic waste disposed of in industrial waste landfills that would result  
29 in CH<sub>4</sub> emissions consists of waste from the pulp and paper and food processing sectors. However, because waste  
30 generation and disposal data are not available in an existing data source for all U.S. industrial waste landfills, a  
31 straight disposal factor is applied over the entire time series to the amount produced to determine the amounts  
32 disposed. Industrial waste facilities reporting under EPA's GHGRP do report detailed waste stream information, and  
33 these data have been used to improve, for example, the DOC value used in the Inventory methodology for the pulp  
34 and paper sector. A 10 percent oxidation factor is also applied to CH<sub>4</sub> generation estimates for industrial waste  
35 landfills, and carries the same amount of uncertainty as with the factor applied to CH<sub>4</sub> generation for MSW landfills.

36 The results of the 2006 IPCC Guidelines Approach 2 quantitative uncertainty analysis are summarized in Table 7-5.  
37 There is considerable uncertainty for the MSW landfills estimates due to the many data sources used, each with its  
38 own uncertainty factor.

**Table 7-5: Approach 2 Quantitative Uncertainty Estimates for CH<sub>4</sub> Emissions from Landfills (MMT CO<sub>2</sub> Eq. and Percent)**

Source	Gas	2017 Emission Estimate (MMT CO <sub>2</sub> Eq.)	Uncertainty Range Relative to Emission Estimate <sup>a</sup>			
			(MMT CO <sub>2</sub> Eq.)		(%)	
			Lower Bound	Upper Bound	Lower Bound	Upper Bound
<b>Total Landfills</b>	<b>CH<sub>4</sub></b>	<b>107.7</b>	<b>95.7</b>	<b>151.2</b>	<b>-11%</b>	<b>40%</b>
MSW	CH <sub>4</sub>	92.8	69.4	116.5	-25%	26%
Industrial	CH <sub>4</sub>	15.0	21.4	41.2	-43%	175%

<sup>a</sup> Range of emission estimates predicted by Monte Carlo Stochastic Simulation for a 95 percent confidence interval.

## QA/QC and Verification

General quality assurance/quality control (QA/QC) procedures were applied consistent with the U.S. QA/QC plan, which is in accordance with Vol. 1 Chapter 6 of *2006 IPCC Guidelines* (see Annex 8 for more details). QA/QC checks are performed for the transcription of the published data set (e.g., EPA’s GHGRP dataset) used to populate the Inventory data set in terms of completeness and accuracy against the reference source. Additionally, all datasets used for this category have been checked to ensure they are of appropriate quality and are representative of U.S. conditions. The primary calculation spreadsheet is tailored from the *2006 IPCC Guidelines* waste model and has been verified previously using the original, peer-reviewed IPCC waste model. All model input values and calculations were verified by secondary QA/QC review. Stakeholder engagements sessions in 2016 and 2017 were used to gather input on methodological improvements and facilitate an external expert review on the methodology, activity data, and emission factors.

Category-specific checks include the following:

- Evaluation of the secondary data sources used as inputs to the Inventory dataset to ensure they are appropriately collected and are reliable;
- Cross-checking the data (activity data and emissions estimates) with previous years to ensure the data are reasonable, and that any significant variation can be explained through the activity data;
- Conducting literature reviews to evaluate the appropriateness of country-specific emission factors (e.g., DOC values, precipitation zones with respect to the application of the k values) given findings from recent peer-reviewed studies; and
- Reviewing secondary datasets to ensure they are nationally complete and supplementing where necessary (e.g., using a scale-up factor to account for emissions from landfills that do not report to EPA’s GHGRP).

A primary focus of the QA/QC checks in past Inventories was to ensure that CH<sub>4</sub> recovery estimates were not double-counted and that all LFGE projects and flares were included in the respective project databases. QA/QC checks performed in the past for the recovery databases were not performed in this Inventory, because new data were not added to the recovery databases in this Inventory year. For the GHGRP data, EPA verifies annual facility-level reports through a multi-step process (e.g., combination of electronic checks and manual reviews by staff) to identify potential errors and ensure that data submitted to EPA are accurate, complete, and consistent. Based on the results of the verification process, EPA follows up with facilities to resolve mistakes that may have occurred.<sup>5</sup>

## Recalculations Discussion

Revisions to the individual facility reports submitted to EPA’s GHGRP can be made at any time and a portion of facilities have revised their reports since 2010 for various reasons, resulting in changes to the total net CH<sub>4</sub> emissions for MSW landfills. These recalculations increased net emissions for MSW landfills from 2005 to 2015 by

<sup>5</sup> See <[https://www.epa.gov/sites/production/files/2015-07/documents/ghgrp\\_verification\\_factsheet.pdf](https://www.epa.gov/sites/production/files/2015-07/documents/ghgrp_verification_factsheet.pdf)>.

1 less than 0.5 percent when compared to the previous Inventory report. Each Inventory year, the back-casted  
2 emissions for 2005 to 2009 will be recalculated using the most recently verified data from the GHGRP. Changes in  
3 these data result in changes to the back-casted emissions.

## 4 **Planned Improvements**

5 EPA has engaged in stakeholder outreach through a series of webinars between December 2016 and August 2017 to  
6 increase the transparency in the Inventory methodology and to identify ideas and supplemental data sources that can  
7 lead to methodological improvements. The areas where EPA is actively working on improvements include the  
8 oxidation factor for 1990 to 2004, the default DOC value, the decay rate (k value), and the scale-up factor.

9 EPA investigated options to adjust the oxidation factor from the 10 percent currently used for 1990 to 2004 to  
10 another value or approach such as the binned approach used in the GHGRP (e.g., 10 percent, 25 percent, or 35  
11 percent based on methane flux). The oxidation factor currently applied in the later portion of the time series (2005 to  
12 2016) averages at 19.5 percent due to the use of the GHGRP data while the earlier portion of the time series applies  
13 the default of 10 percent. No changes to the oxidation factor have been made to the Inventory as a result of EPA's  
14 recent investigations. Efforts will continue to review new literature and revise the value, as appropriate.

15 The Inventory currently uses one value of 0.20 for the DOC for years 1990 to 2004. With respect to improvements  
16 to the DOC value, EPA developed a database with MSW characterization data from individual studies across the  
17 United States. EPA will review this data against the Inventory time series to assess the validity of the current DOC  
18 value and how it is applied in the FOD method. Waste characterization studies vary greatly in terms of the  
19 granularity of waste types included and the spatial boundaries of each study (e.g., one landfill, a metro area,  
20 statewide). EPA also notes longer term recommendation from industry stakeholders regarding the DOC values used  
21 in the GHGRP, in the context of new information on the composition of waste disposed in MSW landfills; these  
22 newer values could then be reflected in the 2005 and later years of the Inventory. EPA is continuing to investigate  
23 publicly available waste characterization studies and calculated DOC values resulting from the study data.

24 EPA began investigating the k values for the three climate types (dry, moderate, and wet) against new data and other  
25 landfill gas models, and how they are applied to the percentage of the population assigned to these climate types.  
26 EPA will also assess the uncertainty factor applied to these k values in the Waste Model. Like the DOC value, the k  
27 values applied through the Waste Model are for the years 1990 to 2004; the k values for 2005 to 2017 are directly  
28 incorporated into the net methane emissions reported to EPA's GHGRP. EPA will continue investigating the  
29 literature for available k value data to understand if the data warrant revisions to the k values used in the Waste  
30 Model between 1990 to 2004.

31 With respect to the scale-up factor, EPA will periodically assess the impact to the waste-in-place and emissions data  
32 from facilities that have resubmitted annual reports during any reporting years, are new reporting facilities, and from  
33 facilities that have stopped reporting to the GHGRP to ensure national estimates are as complete as possible.  
34 Facilities may stop reporting to the GHGRP when they meet the "off-ramp" provisions (reported less than 15,000  
35 metric tons of CO<sub>2</sub> equivalent for 3 consecutive years or less than 25,000 metric tons of CO<sub>2</sub> equivalent for 5  
36 consecutive years). If warranted, EPA will revise the scale-up factor to reflect newly acquired information to ensure  
37 completeness of the Inventory.

38 EPA also conducted a brief investigation of the destruction efficiency applied for landfill gas flares and the  
39 fluctuation in natural gas pricing and other potential factors that are impacting the development of new LFGTE  
40 projects. EPA found that flare destruction efficiencies reported by several vendors ranged from 98 to 99.6 percent.  
41 The EPA applies a 99 percent destruction efficiency for all landfill flares incorporated into the Inventory (from 1990  
42 to 2004 because of the GHGRP data used in later years), which aligns well with the identified range. Therefore, no  
43 revisions have been made to the flare destruction efficiency applied in the Inventory.

### 44 **Box 7-3: Nationwide Municipal Solid Waste Data Sources**

45 Municipal solid waste generated in the United States can be managed through landfilling, recycling, composting,  
46 and combustion with energy recovery. There are three main sources for nationwide solid waste management data in  
47 the United States:

- 48 • The *BioCycle* and Earth Engineering Center of Columbia University's SOG in America surveys [no longer  
49 published];

- The EPA’s *Advancing Sustainable Materials Management: Facts and Figures* reports; and
- The EREF’s *MSW Generation in the United States* reports.

The SOG surveys and, now EREF, collected state-reported data on the amount of waste generated and the amount of waste managed via different management options: landfilling, recycling, composting, and combustion. The survey asked for actual tonnages instead of percentages in each waste category (e.g., residential, commercial, industrial, construction and demolition, organics, tires) for each waste management option. If such a breakdown is not available, the survey asked for total tons landfilled. The data are adjusted for imports and exports across state lines so that the principles of mass balance are adhered to, whereby the amount of waste managed does not exceed the amount of waste generated. The SOG and EREF reports present survey data aggregated to the state level.

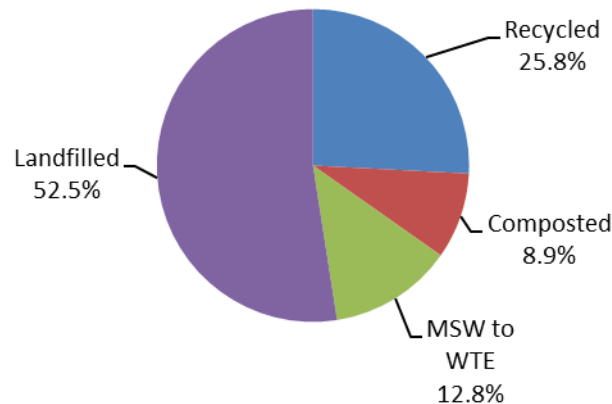
The EPA *Advancing Sustainable Materials Management: Facts and Figures* reports use a materials flow methodology, which relies heavily on a mass balance approach. Data are gathered from industry associations, key businesses, similar industry sources, and government agencies (e.g., the Department of Commerce and the U.S. Census Bureau) and are used to estimate tons of materials and products generated, recycled, combusted with energy recovery or landfilled nationwide. The amount of MSW generated is estimated by estimating production and then adjusting these values by addressing the imports and exports of produced materials to other countries. MSW that is not recycled, composted, or combusted is assumed to be landfilled. The data presented in the report are nationwide totals.

In this Inventory, emissions from solid waste management are presented separately by waste management option, except for recycling of waste materials. Emissions from recycling are attributed to the stationary combustion of fossil fuels that may be used to power on-site recycling machinery, and are presented in the stationary combustion chapter in the Energy sector, although the emissions estimates are not called out separately. Emissions from solid waste disposal in landfills and the composting of solid waste materials are presented in the Landfills and Composting sections in the Waste sector of this report. In the United States, almost all incineration of MSW occurs at waste-to-energy (WTE) facilities or industrial facilities where useful energy is recovered, and thus emissions from waste incineration are accounted for in the Incineration chapter of the Energy sector of this report.

#### Box 7-4: Overview of the Waste Sector

As shown in Figure 7-2 and Figure 7-3, landfilling of MSW is currently and has been the most common waste management practice. A large portion of materials in the waste stream are recovered for recycling and composting, which is becoming an increasingly prevalent trend throughout the country. Materials that are composted and recycled would have previously been disposed in a landfill.

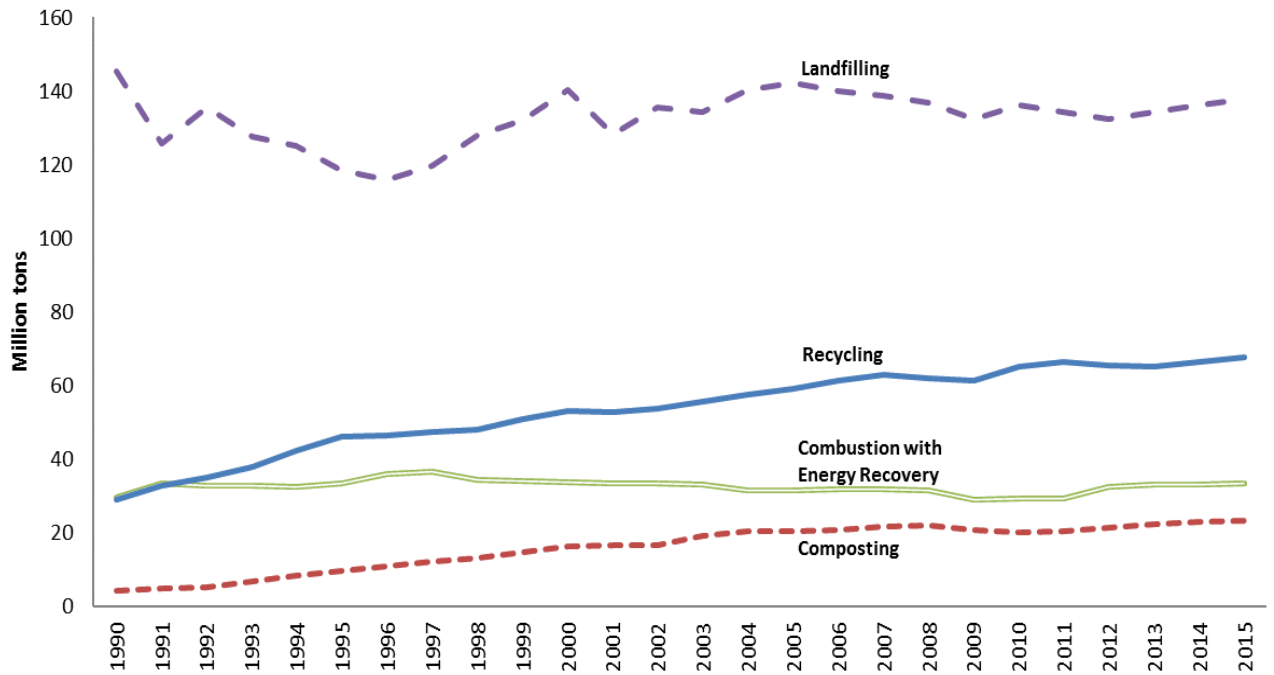
**Figure 7-2: Management of Municipal Solid Waste in the United States, 2015**



Source: EPA (2018c) Note: 2015 is the latest year of available data.



1 **Figure 7-3: MSW Management Trends from 1990 to 2015**



2  
3 Source: EPA (2018c). Note: 2015 is the latest year of available data.

4 Table 7-6 presents a typical composition of waste disposed of at a typical MSW landfill in the United States over  
5 time. It is important to note that the actual composition of waste entering each landfill will vary from that presented  
6 in Table 7-6. Understanding how the waste composition changes over time, specifically for the degradable waste  
7 types (i.e., those types known to generate CH<sub>4</sub> as they break down in a modern MSW landfill), is important for  
8 estimating greenhouse gas emissions. Increased diversion of degradable materials so that they are not disposed of in  
9 landfills reduces the CH<sub>4</sub> generation potential and CH<sub>4</sub> emissions from landfills. For certain degradable waste types  
10 (i.e., paper and paperboard), the amounts discarded have decreased over time due to an increase in waste diversion  
11 through recycling and composting (see Table 7-6 and Figure 7-4). As shown in Figure 7-4, the diversion of food  
12 scraps has been consistently low since 1990 because most cities and counties do not practice curbside collection of  
13 these materials. Neither Table 7-6 nor Figure 7-4 reflect the frequency of backyard composting of yard trimmings  
14 and food waste because this information is largely not collected nationwide and is hard to estimate.

15 **Table 7-6: Materials Discarded<sup>a</sup> in the Municipal Waste Stream by Waste Type from 1990 to**  
16 **2015 (Percent)<sup>b</sup>**

Waste Type	1990	2005	2010	2011 <sup>c</sup>	2012	2013	2014	2015
Paper and Paperboard	30.0%	24.7%	16.1%	14.7%	14.7%	15.0%	14.3%	13.3%
Glass	6.0%	5.8%	5.1%	5.1%	5.2%	5.2%	5.2%	5.1%
Metals	7.2%	7.9%	9.0%	8.9%	9.2%	9.5%	9.5%	9.5%
Plastics	9.5%	16.4%	17.9%	17.9%	18.2%	18.4%	18.5%	18.9%
Rubber and Leather	3.2%	2.9%	3.2%	3.8%	3.2%	3.1%	3.0%	3.3%
Textiles	2.9%	5.3%	6.5%	6.8%	7.1%	7.4%	7.3%	7.6%
Wood	6.9%	7.5%	8.2%	8.2%	8.2%	8.0%	8.1%	8.0%
Other <sup>d</sup>	1.4%	1.8%	2.1%	2.0%	2.0%	1.9%	2.2%	2.2%
Food Scraps	13.6%	18.5%	21.0%	21.4%	21.0%	21.0%	21.7%	22.0%
Yard Trimmings	17.6%	7.0%	8.6%	8.8%	8.7%	8.1%	7.9%	7.8%
Miscellaneous Inorganic Wastes	1.7%	2.2%	2.3%	2.4%	2.4%	2.4%	2.3%	2.3%

<sup>a</sup> Discards after materials and compost recovery. In this table, discards include combustion with energy recovery. Does not include construction & demolition debris, industrial process wastes, or certain other wastes.

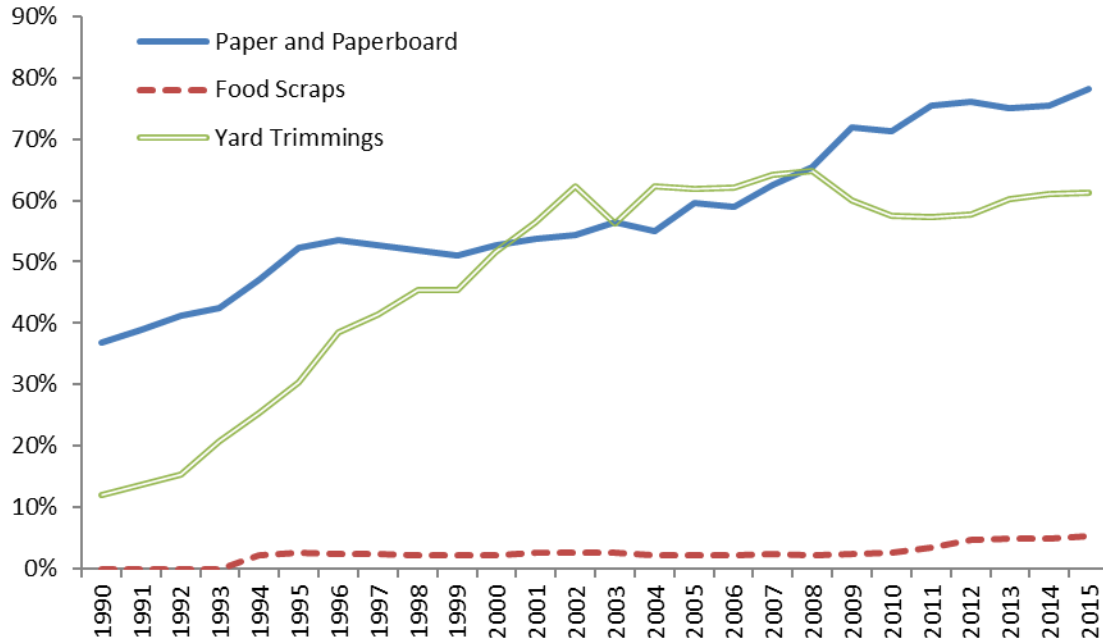
<sup>b</sup> Data for all years except 2011 are from the EPA's *Advancing Sustainable Materials Management: Facts and Figures 2015 Tables and Figures* report (Table 4) published in July 2018 (EPA 2018c).

<sup>c</sup> 2011 data are not included in the most recent *Advancing Sustainable Materials Management: Facts and Figures* report (2014), thus data from the 2013 report (Table 3) was kept in place for 2011 (EPA 2015b).

<sup>d</sup> Includes electrolytes in batteries and fluff pulp, feces, and urine in disposable diapers. Details may not add to totals due to rounding.

Note: 2015 is the latest year of available data.

1 **Figure 7-4: Percent of Degradable Materials Diverted from Landfills from 1990 to 2015**  
2 **(Percent)**



3  
4 Source: (EPA 2018c). Note: 2015 is the latest year of available data.

#### 5 6 **Box 7-5: Description of a Modern, Managed Landfill**

7 Modern, managed landfills are well-engineered facilities that are located, designed, operated, and monitored to  
8 ensure compliance with federal, state, and tribal regulations. Municipal solid waste (MSW) landfills must be  
9 designed to protect the environment from contaminants which may be present in the solid waste stream.  
10 Additionally, many new landfills collect and destroy landfill gas through flares or landfill gas-to-energy projects.  
11 Requirements for affected MSW landfills may include:

- 12 • Siting requirements to protect sensitive areas (e.g., airports, floodplains, wetlands, fault areas, seismic  
13 impact zones, and unstable areas);
- 14 • Design requirements for new landfills to ensure that Maximum Contaminant Levels (MCLs) will not be  
15 exceeded in the uppermost aquifer (e.g., composite liners and leachate collection systems);
- 16 • Leachate collection and removal systems;
- 17 • Operating practices (e.g., daily and intermediate cover, receipt of regulated hazardous wastes, use of  
18 landfill cover material, access options to prevent illegal dumping, use of a collection system to prevent  
19 stormwater run-on/run-off, record-keeping);
- 20 • Air monitoring requirements (explosive gases);
- 21 • Groundwater monitoring requirements;
- 22 • Closure and post-closure care requirements (e.g., final cover construction); and

- Corrective action provisions.

Specific federal regulations that affected MSW landfills must comply with include the 40 CFR Part 258 (Subtitle D of RCRA), or equivalent state regulations and the NSPS 40 CFR Part 60 Subpart W. Additionally, state and tribal requirements may exist.<sup>6</sup>

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## 7.2 Wastewater Treatment (CRF Source Category 5D)

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Wastewater treatment processes can produce anthropogenic methane (CH<sub>4</sub>) and nitrous oxide (N<sub>2</sub>O) emissions. Wastewater from domestic and industrial sources is treated to remove soluble organic matter, suspended solids, pathogenic organisms, and chemical contaminants.<sup>7</sup> Treatment may either occur on site, most commonly through septic systems or package plants, or off site at centralized treatment systems. In the United States, approximately 19 percent of domestic wastewater is treated in septic systems or other on-site systems, while the rest is collected and treated centrally (U.S. Census Bureau 2015). Centralized wastewater treatment systems may include a variety of processes, ranging from lagooning to advanced tertiary treatment technology for removing nutrients. Some wastewater may also be treated through the use of constructed (or semi-natural) wetland systems, though it is much less common in the United States (ERG 2016). Constructed wetlands may be used as the primary method of wastewater treatment, or as a tertiary treatment step following settling and biological treatment. Constructed wetlands develop natural processes that involve vegetation, soil, and associated microbial assemblages to trap and treat incoming contaminants (IPCC 2014).

Soluble organic matter is generally removed using biological processes in which microorganisms consume the organic matter for maintenance and growth. The resulting biomass (sludge) is removed from the effluent prior to discharge to the receiving stream. Microorganisms can biodegrade soluble organic material in wastewater under aerobic or anaerobic conditions, where the latter condition produces CH<sub>4</sub>. During collection and treatment, wastewater may be accidentally or deliberately managed under anaerobic conditions. In addition, the sludge may be further biodegraded under aerobic or anaerobic conditions. The generation of N<sub>2</sub>O may also result from the treatment of domestic wastewater during both nitrification and denitrification of the nitrogen (N) present, usually in the form of urea, ammonia, and proteins. These compounds are converted to nitrate (NO<sub>3</sub>) through the aerobic process of nitrification. Denitrification occurs under anoxic conditions (without free oxygen) and involves the biological conversion of nitrate into dinitrogen gas (N<sub>2</sub>). Nitrous oxide can be an intermediate product of both processes but has typically been associated with denitrification. Recent research suggests that higher emissions of N<sub>2</sub>O may in fact originate from nitrification (Ahn et al. 2010). Other more recent research suggests that N<sub>2</sub>O may also result from other types of wastewater treatment operations (Chandran 2012).

The principal factor in determining the CH<sub>4</sub> generation potential of wastewater is the amount of degradable organic material in the wastewater. Common parameters used to measure the organic component of the wastewater are the biochemical oxygen demand (BOD) and chemical oxygen demand (COD). Under the same conditions, wastewater with higher COD (or BOD) concentrations will generally yield more CH<sub>4</sub> than wastewater with lower COD (or BOD) concentrations. BOD represents the amount of oxygen that would be required to completely consume the organic matter contained in the wastewater through aerobic decomposition processes, while COD measures the total material available for chemical oxidation (both biodegradable and non-biodegradable). The BOD value is most commonly expressed in milligrams of oxygen consumed per liter of sample during 5 days of incubation at 20°C, or BOD<sub>5</sub>. Because BOD is an aerobic parameter, it is preferable to use COD to estimate CH<sub>4</sub> production, since CH<sub>4</sub> is

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<sup>6</sup> For more information regarding federal MSW landfill regulations, see [http://www.epa.gov/osw/nonhaz/municipal/landfill/msw\\_regs.htm](http://www.epa.gov/osw/nonhaz/municipal/landfill/msw_regs.htm).

<sup>7</sup> Throughout the Inventory, emissions from domestic wastewater also include any commercial and industrial wastewater collected and co-treated with domestic wastewater.

1 produced only in anaerobic conditions. The principal factor in determining the N<sub>2</sub>O generation potential of  
 2 wastewater is the amount of N in the wastewater. The variability of N in the influent to the treatment system, as well  
 3 as the operating conditions of the treatment system itself, also impact the N<sub>2</sub>O generation potential.

4 In 2017, CH<sub>4</sub> emissions from domestic wastewater treatment were 8.6 MMT CO<sub>2</sub> Eq. (342 kt CH<sub>4</sub>). Emissions  
 5 remained fairly steady from 1990 through 1999 but have decreased since that time due to decreasing percentages of  
 6 wastewater being treated in anaerobic systems, generally including reduced use of on-site septic systems and central  
 7 anaerobic treatment systems (EPA 1992, 1996, 2000, and 2004; U.S. Census Bureau 2015). In 2017, CH<sub>4</sub> emissions  
 8 from industrial wastewater treatment were estimated to be 5.7 MMT CO<sub>2</sub> Eq. (229 kt CH<sub>4</sub>) and include the newly  
 9 added sector of breweries. Industrial emission sources have generally increased across the time series through 1999  
 10 and then fluctuated up and down with production changes associated with the treatment of wastewater from the pulp  
 11 and paper manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol  
 12 production, petroleum refining, and brewery industries. Table 7-7 and Table 7-8 provide CH<sub>4</sub> emission estimates  
 13 from domestic and industrial wastewater treatment.

14 With respect to N<sub>2</sub>O, the United States identifies two distinct sources for N<sub>2</sub>O emissions from domestic wastewater:  
 15 emissions from centralized wastewater treatment processes, and emissions from effluent from centralized treatment  
 16 systems that has been discharged into aquatic environments. The 2017 emissions of N<sub>2</sub>O from centralized  
 17 wastewater treatment processes and from effluent were estimated to be 0.4 MMT CO<sub>2</sub> Eq. (1.2 kt N<sub>2</sub>O) and 4.6  
 18 MMT CO<sub>2</sub> Eq. (15.4 kt N<sub>2</sub>O), respectively. Total N<sub>2</sub>O emissions from domestic wastewater were estimated to be 5.0  
 19 MMT CO<sub>2</sub> Eq. (16.6 kt N<sub>2</sub>O). Nitrous oxide emissions from wastewater treatment processes gradually increased  
 20 across the time series as a result of increasing U.S. population and protein consumption. Nitrous oxide emissions are  
 21 not estimated from industrial wastewater treatment because there is no IPCC methodology provided or industrial  
 22 wastewater emission factors available. Table 7-7 and Table 7-8 provide N<sub>2</sub>O emission estimates from domestic  
 23 wastewater treatment.

24 **Table 7-7: CH<sub>4</sub> and N<sub>2</sub>O Emissions from Domestic and Industrial Wastewater Treatment**  
 25 **(MMT CO<sub>2</sub> Eq.)**

Activity	1990	2005	2013	2014	2015	2016	2017
<b>CH<sub>4</sub></b>	<b>15.3</b>	<b>15.5</b>	<b>14.4</b>	<b>14.4</b>	<b>14.6</b>	<b>14.3</b>	<b>14.3</b>
Domestic	10.4	10.0	8.9	9.0	9.1	8.7	8.6
Industrial <sup>a</sup>	4.9	5.4	5.5	5.4	5.5	5.6	5.7
<b>N<sub>2</sub>O</b>	<b>3.4</b>	<b>4.4</b>	<b>4.7</b>	<b>4.8</b>	<b>4.8</b>	<b>4.9</b>	<b>5.0</b>
Centralized WWTP	0.2	0.3	0.3	0.3	0.3	0.4	0.4
Domestic Effluent	3.2	4.1	4.3	4.4	4.4	4.5	4.6
<b>Total</b>	<b>18.7</b>	<b>19.8</b>	<b>19.0</b>	<b>19.1</b>	<b>19.3</b>	<b>19.1</b>	<b>19.2</b>

<sup>a</sup> Industrial activity includes the pulp and paper manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol production, petroleum refining, and breweries industries.

Note: Totals may not sum due to independent rounding.

26 **Table 7-8: CH<sub>4</sub> and N<sub>2</sub>O Emissions from Domestic and Industrial Wastewater Treatment (kt)**

Activity	1990	2005	2013	2014	2015	2016	2017
<b>CH<sub>4</sub></b>	<b>612</b>	<b>618</b>	<b>574</b>	<b>575</b>	<b>582</b>	<b>571</b>	<b>571</b>
Domestic	418	401	355	359	363	347	342
Industrial <sup>a</sup>	194	217	219	216	219	224	229
<b>N<sub>2</sub>O</b>	<b>11</b>	<b>15</b>	<b>16</b>	<b>16</b>	<b>16</b>	<b>16</b>	<b>17</b>
Centralized WWTP	1	1	1	1	1	1	1
Domestic Effluent	11	14	15	15	15	15	15

<sup>a</sup> Industrial activity includes pulp and paper manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol production, petroleum refining, and breweries.

Note: Totals may not sum due to independent rounding.

# Methodology

## Domestic Wastewater CH<sub>4</sub> Emission Estimates

Domestic wastewater CH<sub>4</sub> emissions originate from both septic systems and from centralized treatment systems, such as publicly owned treatment works (POTWs). Within these centralized systems, CH<sub>4</sub> emissions can arise from aerobic systems that are not well managed or that are designed to have periods of anaerobic activity (e.g., constructed wetlands and facultative lagoons), anaerobic systems (anaerobic lagoons and anaerobic reactors), and from anaerobic digesters when the captured biogas is not completely combusted. The methodological equations are:

$$\begin{aligned} \text{Emissions from Septic Systems} &= A \\ &= \text{US}_{\text{POP}} \times (\% \text{ onsite}) \times (\text{EF}_{\text{SEPTIC}}) \times 1/10^9 \times 365.25 \end{aligned}$$

$$\begin{aligned} &\text{Emissions from Centrally Treated Aerobic Systems (other than Constructed Wetlands)} + \text{Emissions from} \\ &\text{Centrally Treated Aerobic Systems (Constructed Wetlands Only)} + \text{Emissions from Centrally Treated Aerobic} \\ &\text{Systems (Constructed Wetlands used as Tertiary Treatment)} = B \end{aligned}$$

where,

$$\begin{aligned} &\text{Emissions from Centrally Treated Aerobic Systems (other than Constructed Wetlands)} \\ &= [(\% \text{ collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ aerobic}_{\text{COTCW}}) \times (\% \text{ aerobic w/out primary}) + (\% \text{ collected}) \times \\ &(\text{total BOD}_5 \text{ produced}) \times (\% \text{ aerobic}_{\text{COTCW}}) \times (\% \text{ aerobic w/primary}) \times (1 - \% \text{ BOD removed in prim. treat.})] \times \\ &(\% \text{ operations not well managed}) \times (B_o) \times (\text{MCF-aerobic\_not\_well\_man}) \end{aligned}$$

$$\begin{aligned} &\text{Emissions from Centrally Treated Aerobic Systems (Constructed Wetlands Only)} \\ &= [(\% \text{ collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ aerobic}_{\text{CW}})] \times (B_o) \times (\text{MCF-constructed wetlands}) \end{aligned}$$

$$\begin{aligned} &\text{Emissions from Centrally Treated Aerobic Systems (Constructed Wetlands used as Tertiary Treatment)} \\ &= [(\text{POTW\_flow\_CW}) \times (\text{BOD}_{\text{CW,INF}}) \times 3.79] \times 1/10^6 \times 365.25 \end{aligned}$$

$$\begin{aligned} &\text{Emissions from Centrally Treated Anaerobic Systems} = C \\ &= \{[(\% \text{ collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ anaerobic}) \times (\% \text{ anaerobic w/out primary})] + [(\% \\ &\text{collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ anaerobic}) \times (\% \text{ anaerobic w/primary}) \times (1 - \% \text{ BOD removed in} \\ &\text{prim. treat.})]\} \times (B_o) \times (\text{MCF-anaerobic}) \end{aligned}$$

$$\begin{aligned} &\text{Emissions from Anaerobic Digesters} = D \\ &= [(\text{POTW\_flow\_AD}) \times (\text{digester gas}) / (100)] \times 0.0283 \times (\text{FRAC}_{\text{CH}_4}) \times 365.25 \times (662) \times (1 - \text{DE}) \times 1/10^9 \end{aligned}$$

$$\text{Total Domestic CH}_4 \text{ Emissions from Wastewater (kt)} = A + B + C + D$$

where,

US <sub>POP</sub>	= U.S. population
% onsite	= Flow to septic systems / total flow
% collected	= Flow to POTWs / total flow
% aerobic <sub>COTCW</sub>	= Flow to aerobic systems, other than wetlands only / total flow to POTWs
% aerobic <sub>CW</sub>	= Flow to aerobic systems, constructed wetlands used as sole treatment / total flow to POTWs
% anaerobic	= Flow to anaerobic systems / total flow to POTWs
% aerobic w/out primary	= Percent of aerobic systems that do not employ primary treatment
% aerobic w/primary	= Percent of aerobic systems that employ primary treatment
% BOD removed in prim. treat.	= Percent of BOD removed in primary treatment
% operations not well managed	= Percent of aerobic systems that are not well managed and in which some anaerobic degradation occurs
% anaerobic w/out primary	= Percent of anaerobic systems that do not employ primary treatment
% anaerobic w/primary	= Percent of anaerobic systems that employ primary treatment

1	EF <sub>SEPTIC</sub>	= Methane emission factor – septic systems
2	Total BOD <sub>5</sub> produced	= kg BOD/capita/day × U.S. population × 365.25 days/yr
3	BOD <sub>CW,INF</sub>	= BOD concentration in wastewater entering the constructed wetland
4	B <sub>o</sub>	= Maximum CH <sub>4</sub> -producing capacity for domestic wastewater
5	1/10 <sup>6</sup>	= Conversion factor, kg to kt
6	365.25	= Days in a year
7	3.79	= Conversion factor, gallons to liters
8	MCF-aerobic_not_well_man.	= CH <sub>4</sub> correction factor for aerobic systems that are not well managed
9	MCF-anaerobic	= CH <sub>4</sub> correction factor for anaerobic systems
10	MCF-constructed wetlands	= CH <sub>4</sub> correction factor for surface flow constructed wetlands
11	DE	= CH <sub>4</sub> destruction efficiency from flaring or burning in engine
12	POTW_flow_CW	= Wastewater flow to POTWs that use constructed wetlands as tertiary treatment (MGD)
13		
14	POTW_flow_AD	= Wastewater influent flow to POTWs that have anaerobic digesters (MGD)
15		
16	digester gas	= Cubic feet of digester gas produced per person per day
17	100	= Wastewater flow to POTW (gallons/person/day)
18	0.0283	= Conversion factor, ft <sup>3</sup> to m <sup>3</sup>
19	FRAC_CH <sub>4</sub>	= Proportion of CH <sub>4</sub> in biogas
20	662	= Density of CH <sub>4</sub> (g CH <sub>4</sub> /m <sup>3</sup> CH <sub>4</sub> )
21	1/10 <sup>9</sup>	= Conversion factor, g to kt

22 **Emissions from Septic Systems:**

23 Methane emissions from septic systems were estimated by multiplying the U.S. population by the percent of  
 24 wastewater treated in septic systems (about 18 percent) and an emission factor (10.7 g CH<sub>4</sub>/capita/day) (Leverenz et  
 25 al. 2010), and then converting the result to kt/year. U.S. population data were taken from the U.S. Census Bureau  
 26 International Database (U.S. Census Bureau 2018) and include the populations of the United States, American  
 27 Samoa, Guam, Northern Mariana Islands, Puerto Rico, and the U.S. Virgin Islands. Table 7-9 presents U.S.  
 28 population for 1990 through 2017.

29 **Emissions from Centrally Treated Aerobic and Anaerobic Systems:**

30 Methane emissions from POTWs were estimated by multiplying the total BOD<sub>5</sub> produced in the United States by the  
 31 percent of wastewater treated centrally, or percent collected (about 82 percent) (U.S. Census Bureau 2015), the  
 32 relative percentage of wastewater treated by aerobic and anaerobic systems (other than constructed wetlands), the  
 33 relative percentage of aerobic systems at wastewater facilities with and without primary treatment (EPA 1992, 1996,  
 34 2000, and 2004), the relative percentage of anaerobic systems at wastewater facilities with and without primary  
 35 treatment (EPA 1992, 1996, 2000, and 2004), the percentage of BOD<sub>5</sub> treated after primary treatment (67.5 percent,  
 36 32.5 percent removed in primary treatment) (Metcalf & Eddy 2014), the maximum CH<sub>4</sub>-producing capacity of  
 37 domestic wastewater (B<sub>o</sub>, 0.6 kg CH<sub>4</sub>/kg BOD) (IPCC 2006), and the relative methane correction factors (MCF) not  
 38 well managed aerobic (0.3) (IPCC 2006), and anaerobic (0.8) (IPCC 2006) systems. All aerobic systems are  
 39 assumed to be well-managed as there are currently no data available to quantify the number of systems that are not  
 40 well-managed.

41 Table 7-9 presents total BOD<sub>5</sub> produced for 1990 through 2017. The proportions of domestic wastewater treated  
 42 onsite versus at centralized treatment plants were based on data from the 1989, 1991, 1993, 1995, 1997, 1999, 2001,  
 43 2003, 2005, 2007, 2009, 2011, 2013, and 2015 *American Housing Surveys* conducted by the U.S. Census Bureau  
 44 (U.S. Census Bureau 2015), with data for intervening years obtained by linear interpolation and 2017 forecasted  
 45 using 1990 to 2016 data. The BOD<sub>5</sub> production rate was determined using BOD generation rates per capita both  
 46 with and without kitchen scraps (Metcalf & Eddy 2003; Metcalf & Eddy 2014) as well as an estimated percent of  
 47 housing units that utilize kitchen garbage disposals (ERG 2018a). The percent BOD<sub>5</sub> removed by primary treatment  
 48 for domestic wastewater was obtained from Metcalf & Eddy (2014). The percent of wastewater flow to aerobic and  
 49 anaerobic systems, the percent of aerobic and anaerobic systems that do and do not employ primary treatment, and  
 50 the wastewater flow to POTWs that have anaerobic digesters were obtained from the 1992, 1996, 2000, and 2004  
 51 Clean Watersheds Needs Survey (EPA 1992, 1996, 2000, and 2004). Data for intervening years were obtained by  
 52 linear interpolation and the years 2005 through 2016 were forecasted from the rest of the time series. The percent of  
 53 wastewater flow to aerobic systems that use only constructed wetlands and wastewater flow to POTWs that use

1 constructed wetlands as tertiary treatment were obtained from the 1992, 1996, 2000, 2004, 2008, and 2012 Clean  
 2 Watersheds Needs Survey (EPA 1992, 1996, 2000, 2004, 2008b, and 2012). Data for intervening years were  
 3 obtained by linear interpolation and the years 2013 through 2017 were forecasted from the rest of the time series.

4 **Table 7-9: U.S. Population (Millions) and Domestic Wastewater BOD<sub>5</sub> Produced (kt)**

Year	Population	BOD <sub>5</sub>
1990	253	8,131
2005	300	9,624
2013	320	9,672
2014	323	9,656
2015	325	9,739
2016	327	9,820
2017	330	9,938

Sources: U.S. Census Bureau (2018); ERG (2018a).

5 For constructed wetlands, an MCF of 0.4 was used, which is the IPCC suggested MCF for surface flow wetlands.  
 6 This is the most conservative factor for constructed wetlands and was recommended by IPCC (2014) when the type  
 7 of constructed wetland is not known. A BOD<sub>5</sub> concentration of 30 mg/L was used for wastewater entering  
 8 constructed wetlands used as tertiary treatment based on U.S. secondary treatment standards for POTWs. These  
 9 standards are based on plants generally utilizing simple settling and biological treatment (EPA 2013).

10 In addition, methane emissions were calculated for systems that treat wastewater with constructed wetlands and  
 11 systems that use constructed wetlands as tertiary treatment; however, constructed wetlands are a relatively small  
 12 portion of wastewater treated centrally (<0.1 percent).

13 **Emissions from Anaerobic Digesters:**

14 Total CH<sub>4</sub> emissions from anaerobic digesters were estimated by multiplying the wastewater influent flow to  
 15 POTWs with anaerobic digesters, the cubic feet of digester gas generated per person per day divided by the flow to  
 16 POTWs, the fraction of CH<sub>4</sub> in biogas (0.65), the density of CH<sub>4</sub> (662 g CH<sub>4</sub>/m<sup>3</sup> CH<sub>4</sub>) (EPA 1993a), one minus the  
 17 destruction efficiency from burning the biogas in an energy/thermal device (0.99 for enclosed flares) and then  
 18 converting the results to kt/year.

19 The CH<sub>4</sub> destruction efficiency for CH<sub>4</sub> recovered from sludge digestion operations, 99 percent, was selected based  
 20 on the range of efficiencies (98 to 100 percent) recommended for flares in *AP-42 Compilation of Air Pollutant*  
 21 *Emission Factors*, Chapter 2.4 (EPA 1998), along with data from CAR (2011), Sullivan (2007), Sullivan (2010), and  
 22 UNFCCC (2012). The cubic feet of digester gas produced per person per day (1.0 ft<sup>3</sup>/person/day) and the proportion  
 23 of CH<sub>4</sub> in biogas (0.65) come from Metcalf & Eddy (2014). The wastewater flow to a POTW (100 gal/person/day)  
 24 was taken from the Great Lakes-Upper Mississippi River Board of State and Provincial Public Health and  
 25 Environmental Managers, "*Recommended Standards for Wastewater Facilities (Ten-State Standards)*" (2004).

26 Table 7-10 presents domestic wastewater CH<sub>4</sub> emissions for both septic and centralized systems, including  
 27 anaerobic digesters, in 2017.

28 **Table 7-10: Domestic Wastewater CH<sub>4</sub> Emissions from Septic and Centralized Systems**  
 29 **(2017, MMT CO<sub>2</sub> Eq. and Percent)**

	CH <sub>4</sub> Emissions (MMT CO <sub>2</sub> Eq.)	% of Domestic Wastewater CH <sub>4</sub>
Septic Systems	5.9	69.1%
Centrally-Treated Aerobic Systems	0.1	1.2%
Centrally-Treated Anaerobic Systems	2.3	27.3%
Anaerobic Digesters	0.2	2.4%
<b>Total</b>	<b>8.6</b>	<b>100%</b>

Note: Totals may not sum due to independent rounding.

## 1 Industrial Wastewater CH<sub>4</sub> Emission Estimates

2 Methane emission estimates from industrial wastewater were developed according to the methodology described in  
 3 the *2006 IPCC Guidelines*. Industry categories that are likely to produce significant CH<sub>4</sub> emissions from wastewater  
 4 treatment were identified and included in the Inventory. The main criteria used to identify these industries are  
 5 whether they generate high volumes of wastewater, whether there is a high organic wastewater load, and whether the  
 6 wastewater is treated using methods that result in CH<sub>4</sub> emissions. The top six industries that meet these criteria are  
 7 pulp and paper manufacturing; meat and poultry processing; vegetables, fruits, and juices processing; starch-based  
 8 ethanol production; petroleum refining; and breweries. Wastewater treatment emissions for these sectors for 2017  
 9 are displayed in Table 7-11 below. Table 7-12 contains production data for these industries.

10 **Table 7-11: Industrial Wastewater CH<sub>4</sub> Emissions by Sector (2017, MMT CO<sub>2</sub> Eq. and**  
 11 **Percent)**

	CH <sub>4</sub> Emissions (MMT CO <sub>2</sub> Eq.)	% of Industrial Wastewater CH <sub>4</sub>
Meat & Poultry	4.7	81.5%
Pulp & Paper	0.6	10.0%
Fruit & Vegetables	0.1	2.4%
Petroleum Refineries	0.1	2.6%
Ethanol Refineries	0.1	2.6%
Breweries	0.05	1%
<b>Total</b>	<b>5.7</b>	<b>100%</b>

Note: Totals may not sum due to independent rounding.

12 **Table 7-12: U.S. Pulp and Paper, Meat, Poultry, Vegetables, Fruits and Juices, Ethanol,**  
 13 **Breweries, and Petroleum Refining Production (MMT)**

Year	Pulp and Paper <sup>a</sup>	Meat (Live Weight Killed)	Poultry (Live Weight Killed)	Vegetables, Fruits and Juices	Ethanol	Breweries	Petroleum Refining
1990	82.5	27.3	14.6	38.7	2.5	23.9	702.4
2005	91.8	31.4	25.1	42.9	11.7	23.2	818.6
2013	79.9	33.6	26.5	45.1	39.7	22.5	878.7
2014	80.9	32.2	26.9	45.3	42.8	22.5	903.9
2015	80.9	32.8	27.7	44.6	44.2	22.4	914.5
2016	79.9	34.2	28.3	43.2	45.8	22.3	926.0
2017	79.9	35.4	28.9	42.60	47.2	21.8	934.1

<sup>a</sup>Pulp and paper production is the sum of market pulp production plus paper and paperboard production.

Sources: FAO (2018a) and FAO (2018b); USDA (2018a); Cooper (2018); Beer Institute (2011) and TTB (2018); EIA (2018).

14 Methane emissions from these categories were estimated by multiplying the annual product output by the average  
 15 outflow, the organics loading (in COD) in the outflow, the maximum CH<sub>4</sub> producing potential of industrial  
 16 wastewater (B<sub>0</sub>), and the percentage of organic loading assumed to degrade anaerobically in a given treatment  
 17 system (MCF). Ratios of BOD:COD in various industrial wastewaters were obtained from EPA (1997a) and used to  
 18 estimate COD loadings. The B<sub>0</sub> value used for all industries is the IPCC default value of 0.25 kg CH<sub>4</sub>/kg COD  
 19 (IPCC 2006).

20 For each industry, the percent of plants in the industry that treat wastewater on site, the percent of plants that have a  
 21 primary treatment step prior to biological treatment, and the percent of plants that treat wastewater anaerobically  
 22 were defined. The percent of wastewater treated anaerobically onsite (TA) was estimated for both primary treatment  
 23 (%TA<sub>p</sub>) and secondary treatment (%TA<sub>s</sub>). For plants that have primary treatment in place, an estimate of COD that  
 24 is removed prior to wastewater treatment in the anaerobic treatment units was incorporated. The values used in the  
 25 %TA calculations are presented in Table 7-13 below.



1 The methodological equations are:

$$2 \quad \text{CH}_4 (\text{industrial wastewater}) = [P \times W \times \text{COD} \times \%TA_p \times B_o \times \text{MCF}] + [P \times W \times \text{COD} \times \%TA_s \times B_o \times \text{MCF}]$$

$$3 \quad \%TA_p = [\%Plants_o \times \%WW_{a,p} \times \%COD_p]$$

$$4 \quad \%TA_s = [\%Plants_a \times \%WW_{a,s} \times \%COD_s] + [\%Plants_t \times \%WW_{a,t} \times \%COD_s]$$

5 where,

6  $\text{CH}_4$  (industrial wastewater) = Total  $\text{CH}_4$  emissions from industrial wastewater (kg/year)

7 P = Industry output (metric tons/year)

8 W = Wastewater generated ( $\text{m}^3$ /metric ton of product)

9 COD = Organics loading in wastewater ( $\text{kg}/\text{m}^3$ )

10  $\%TA_p$  = Percent of wastewater treated anaerobically on site in primary treatment

11  $\%TA_s$  = Percent of wastewater treated anaerobically on site in secondary treatment

12  $\%Plants_o$  = Percent of plants with onsite treatment

13  $\%WW_{a,p}$  = Percent of wastewater treated anaerobically in primary treatment

14  $\%COD_p$  = Percent of COD entering primary treatment

15  $\%Plants_a$  = Percent of plants with anaerobic secondary treatment

16  $\%Plants_t$  = Percent of plants with other secondary treatment

17  $\%WW_{a,s}$  = Percent of wastewater treated anaerobically in anaerobic secondary treatment

18  $\%WW_{a,t}$  = Percent of wastewater treated anaerobically in other secondary treatment

19  $\%COD_s$  = Percent of COD entering secondary treatment

20  $B_o$  = Maximum  $\text{CH}_4$  producing potential of industrial wastewater (kg  $\text{CH}_4$ /kg COD)

21 MCF =  $\text{CH}_4$  correction factor, indicating the extent to which the organic content (measured as COD) degrades anaerobically

24 Alternate methodological equations for calculating %TA were used for secondary treatment in the pulp and paper industry to account for aerobic systems with anaerobic portions. These equations are:

$$26 \quad \%TA_a = [\%Plants_a \times \%WW_{a,s} \times \%COD_s] + [\%Plants_{a,t} \times \%WW_{a,t} \times \%COD_s]$$

$$27 \quad \%TA_{a,t} = [\%Plants_{a,t} \times \%WW_{a,t} \times \%COD_s]$$

28 where,

29  $\%TA_a$  = Percent of wastewater treated anaerobically on site in secondary treatment

30  $\%TA_{a,t}$  = Percent of wastewater treated in aerobic systems with anaerobic portions on site in secondary treatment

31  $\%Plants_a$  = Percent of plants with anaerobic secondary treatment

32  $\%Plants_{a,t}$  = Percent of plants with partially anaerobic secondary treatment

33  $\%WW_{a,s}$  = Percent of wastewater treated anaerobically in anaerobic secondary treatment

34  $\%WW_{a,t}$  = Percent of wastewater treated anaerobically in other secondary treatment

35  $\%COD_s$  = Percent of COD entering secondary treatment

37 As described below, the values presented in Table 7-13 were used in the emission calculations and are described in detail in ERG (2008), ERG (2013a), and ERG (2013b).

39 **Table 7-13: Variables Used to Calculate Percent Wastewater Treated Anaerobically by Industry (Percent)**

Variable	Industry								
	Pulp and Paper	Meat Processing	Poultry Processing	Fruit/Vegetable Processing	Ethanol Production – Wet Mill	Ethanol Production – Dry Mill	Petroleum Refining	Breweries – Craft	Breweries – Non-Craft
$\%TA_p$	0	0	0	0	0	0	0	0	0
$\%TA_s$	0	33	25	4.2	33.3	75	23.6	0	0
$\%TA_a$	2.2	0	0	0	0	0	0	0	0
$\%TA_{a,t}$	11.8	0	0	0	0	0	0	0	0
$\%Plants_o$	0	100	100	11	100	100	100	100	1

%Plants <sub>a</sub>	5	33	25	5.5	33.3	75	23.6	0	0
%Plants <sub>a,t</sub>	28	0	0	0	0	0	0	0	0
%Plants <sub>t</sub>	35	67	75	5.5	66.7	25	0	0	0
%WW <sub>a,p</sub>	0	0	0	0	0	0	0	0	0
%WW <sub>a,s</sub>	100	100	100	100	100	100	100	0	0
%WW <sub>a,t</sub>	0	0	0	0	0	0	0	0	0
%COD <sub>p</sub>	100	100	100	100	100	100	100	0	0
%COD <sub>s</sub>	42	100	100	77	100	100	100	0	0

Note: Due to differences in data availability and methodology, zero values in the table are for calculation purposes only and may indicate unavailable data.

Sources: ERG (2008); ERG (2013a); and ERG (2013b).

1 *Pulp and Paper.* Wastewater treatment for the pulp and paper industry typically includes neutralization, screening,  
2 sedimentation, and flotation/hydrocycloning to remove solids (World Bank 1999; Nemerow and Dasgupta 1991).  
3 Secondary treatment (storage, settling, and biological treatment) mainly consists of lagooning. In determining the  
4 percent that degrades anaerobically, both primary and secondary treatment were considered. In the United States,  
5 primary treatment is focused on solids removal, equalization, neutralization, and color reduction (EPA 1993b). The  
6 vast majority of pulp and paper mills with on-site treatment systems use mechanical clarifiers to remove suspended  
7 solids from the wastewater. About 10 percent of pulp and paper mills with treatment systems use settling ponds for  
8 primary treatment and these are more likely to be located at mills that do not perform secondary treatment (EPA  
9 1993b). However, because the vast majority of primary treatment operations at U.S. pulp and paper mills use  
10 mechanical clarifiers, and less than 10 percent of pulp and paper wastewater is managed in primary settling ponds  
11 that are not expected to have anaerobic conditions, negligible emissions are assumed to occur during primary  
12 treatment.

13 Approximately 42 percent of the BOD passes on to secondary treatment, which consists of activated sludge, aerated  
14 stabilization basins, or non-aerated stabilization basins. Based on EPA's *OAQPS Pulp and Paper Sector Survey*, 5.3  
15 percent of pulp and paper mills reported using anaerobic secondary treatment for wastewater and/or pulp  
16 condensates (ERG 2013a). Twenty-eight percent of mills also reported the use of quiescent settling ponds. Using  
17 engineering judgment, these systems were determined to be aerobic with possible anaerobic portions. For the truly  
18 anaerobic systems, an MCF of 0.8 is used, as these are typically deep stabilization basins. For the partially anaerobic  
19 systems, an MCF of 0.2 is used, which is the *2006 IPCC Guidelines*-suggested MCF for shallow lagoons.

20 A time series of CH<sub>4</sub> emissions for 1990 through 2017 was developed based on paper and paperboard production  
21 data from the Food and Agricultural Organization of the United Nations (FAO) database FAOSTAT. (FAO 2018a)  
22 and market pulp production data from FAO Pulp and Paper Capacities Reports (FAO 2018b). Market pulp  
23 production values were available directly for 1998, 2000 through 2004, and 2010 through 2016. Where market pulp  
24 data were unavailable, a percent of woodpulp that is market pulp was applied to woodpulp production values from  
25 FAOSTAT to estimate market pulp production (FAO 2018a). The percent of woodpulp that is market pulp for 1990-  
26 1997 was assumed to be the same as 1998, 1999 was interpolated between values for 1998 and 2000, 2000 through  
27 2009 were interpolated between values for 2003 and 2010, and 2017 was forecasted from the rest of the time series.  
28 A time series of the overall wastewater outflow for 1990 through 1994 varies based on data outlined in ERG (2013a)  
29 to reflect historical wastewater flow. Wastewater generation rates for 1995, 2000, and 2002 were estimated from the  
30 *2014 American Forest and Paper Association (AF&PA) Sustainability Report* (AF&PA 2014). Wastewater  
31 generation rates for 2004, 2006, 2008, 2010, 2012, and 2014 were estimated from the *2016 AF&PA Sustainability*  
32 *Report* (AF&PA 2016). Data for intervening years were obtained by linear interpolation, while 2015 through 2017  
33 were forecasted from the rest of the time series. The average BOD concentrations in raw wastewater was estimated  
34 to be 0.4 grams BOD/liter for 1990 to 1998, while 0.3 grams BOD/liter was estimated for 2014 through 2017 (EPA  
35 1997b; EPA 1993b; World Bank 1999; Malmberg 2018). Data for intervening years were obtained by linear  
36 interpolation. The COD:BOD ratio used to convert the organic loading to COD for pulp and paper mills was 2.5 for  
37 the entire time series (Malmberg 2018).

38 *Meat and Poultry Processing.* The meat and poultry processing industry makes extensive use of anaerobic lagoons  
39 in sequence with screening, fat traps, and dissolved air flotation when treating wastewater on site. About 33 percent  
40 of meat processing operations (EPA 2002) and 25 percent of poultry processing operations (U.S. Poultry 2006)  
41 perform on-site treatment in anaerobic lagoons. The IPCC default B<sub>0</sub> of 0.25 kg CH<sub>4</sub>/kg COD and default MCF of  
42 0.8 for anaerobic lagoons were used to estimate the CH<sub>4</sub> produced from these on-site treatment systems. Production

1 data on carcass weight and live weight killed for the meat and poultry industry were obtained from the USDA  
 2 *Agricultural Statistics Database and the Agricultural Statistics Annual Reports* (USDA 2018a). Data collected by  
 3 EPA’s Office of Water provided estimates for wastewater flows into anaerobic lagoons: 5.3 and 12.5 m<sup>3</sup>/metric ton  
 4 for meat and poultry production (live weight killed), respectively (EPA 2002). The loadings are 2.8 and 1.5 g  
 5 BOD/liter for meat and poultry, respectively (EPA 2002). The COD:BOD ratio used to convert the organic loading  
 6 to COD for both meat and poultry facilities was 3 (EPA 1997a).

7 *Vegetables, Fruits, and Juices Processing*. Treatment of wastewater from fruits, vegetables, and juices processing  
 8 includes screening, coagulation/settling, and biological treatment (lagooning). The flows are frequently seasonal,  
 9 and robust treatment systems are preferred for on-site treatment. Effluent is suitable for discharge to POTWs. This  
 10 industry is likely to use lagoons intended for aerobic operation, but the large seasonal loadings may develop limited  
 11 anaerobic zones. In addition, some anaerobic lagoons may also be used (Nemerow and Dasgupta 1991).  
 12 Consequently, 4.2 percent of these wastewater organics are assumed to degrade anaerobically (ERG 2008). The  
 13 IPCC default B<sub>0</sub> of 0.25 kg CH<sub>4</sub>/kg COD and default MCF of 0.8 for anaerobic treatment were used to estimate the  
 14 CH<sub>4</sub> produced from these on-site treatment systems. The USDA National Agricultural Statistics Service (USDA  
 15 2018a, 2018c) provided production data for potatoes, other vegetables, citrus fruit, non-citrus fruit, and grapes  
 16 processed for wine. Outflow and BOD data, presented in Table 7-14, were obtained from EPA (1974) for potato,  
 17 citrus fruit, and apple processing, and from EPA (1975) for all other commodities. The COD:BOD ratio used to  
 18 convert the organic loading to COD for all fruit, vegetable, and juice facilities was 1.5 (EPA 1997a).

19 **Table 7-14: Wastewater Flow (m<sup>3</sup>/ton) and BOD Production (g/L) for U.S. Vegetables,**  
 20 **Fruits, and Juices Production**

Commodity	Wastewater Outflow (m <sup>3</sup> /ton)	BOD (g/L)
<b>Vegetables</b>		
Potatoes	10.27	1.765
Other Vegetables	8.55	0.776
<b>Fruit</b>		
Apples	3.66	1.371
Citrus Fruits	10.11	0.317
Non-citrus Fruits	12.42	1.204
Grapes (for wine)	2.78	1.831

Sources: EPA (1974); EPA (1975).

21 *Ethanol Production*. Ethanol, or ethyl alcohol, is produced primarily for use as a fuel component, but is also used in  
 22 industrial applications and in the manufacture of beverage alcohol. Ethanol can be produced from the fermentation  
 23 of sugar-based feedstocks (e.g., molasses and beets), starch- or grain-based feedstocks (e.g., corn, sorghum, and  
 24 beverage waste), and cellulosic biomass feedstocks (e.g., agricultural wastes, wood, and bagasse). Ethanol can also  
 25 be produced synthetically from ethylene or hydrogen and carbon monoxide. However, synthetic ethanol comprises  
 26 only about 2 percent of ethanol production and is only in an experimental stage in the United States. Currently,  
 27 ethanol is mostly made from sugar and starch crops, but with advances in technology, cellulosic biomass is  
 28 increasingly used as ethanol feedstock (DOE 2013).

29 Ethanol is produced from corn (or other starch-based feedstocks) primarily by two methods: wet milling and dry  
 30 milling. Historically, the majority of ethanol was produced by the wet milling process, but now the majority is  
 31 produced by the dry milling process. The dry milling process is cheaper to implement and has become more efficient  
 32 in recent years (Rendleman and Shapouri 2007). The wastewater generated at ethanol production facilities is  
 33 handled in a variety of ways. Dry milling facilities often combine the resulting evaporator condensate with other  
 34 process wastewaters, such as equipment wash water, scrubber water, and boiler blowdown and anaerobically treat  
 35 this wastewater using various types of digesters. Wet milling facilities often treat their steepwater condensate in  
 36 anaerobic systems followed by aerobic polishing systems. Wet milling facilities may treat the stillage (or processed  
 37 stillage) from the ethanol fermentation/distillation process separately or together with steepwater and/or wash water.  
 38 Methane generated in anaerobic digesters is commonly collected and either flared or used as fuel in the ethanol  
 39 production process (ERG 2006).

40 Available information was compiled from the industry on wastewater generation rates, which ranged from 1.25  
 41 gallons per gallon ethanol produced (for dry milling) to 10 gallons per gallon ethanol produced (for wet milling)

(Ruocco 2006a; Ruocco 2006b; Merrick 1998; Donovan 1996; NRBP 2001). COD concentrations were found to be about 3 g/L (Ruocco 2006a; Merrick 1998; White and Johnson 2003). One hundred percent of plants were estimated to have onsite wastewater treatment, and the variables used to calculate percent wastewater treated anaerobically are presented in Table 7-13. A default MCF of 0.8 for anaerobic treatment was used to estimate the CH<sub>4</sub> produced from these on-site treatment systems. The amount of CH<sub>4</sub> recovered through the use of biomethanators was estimated, and a 99 percent destruction efficiency was used. Biomethanators are anaerobic reactors that use microorganisms under anaerobic conditions to reduce COD and organic acids and recover biogas from wastewater (ERG 2006). Methane emissions for dry milling and wet milling processes were then estimated as follows:

$$\text{Methane} = [\text{Production} \times \text{Flow} \times \text{COD} \times 3.785 \times (\% \text{Plants}_o \times \% \text{WW}_{a,p} \times \% \text{COD}_p) + [\% \text{Plants}_a \times \% \text{WW}_{a,s} \times \% \text{COD}_s] + [\% \text{Plants}_t \times \% \text{WW}_{a,t} \times \% \text{COD}_s]) \times B_o \times \text{MCF} \times \% \text{ Not Recovered}] + [\text{Production} \times \text{Flow} \times 3.785 \times \text{COD} \times (\% \text{Plants}_o \times \% \text{WW}_{a,p} \times \% \text{COD}_p) + [\% \text{Plants}_a \times \% \text{WW}_{a,s} \times \% \text{COD}_s] + [\% \text{Plants}_t \times \% \text{WW}_{a,t} \times \% \text{COD}_s]) \times B_o \times \text{MCF} \times (\% \text{ Recovered}) \times (1 - \text{DE})] \times 1/10^9$$

where,

Production	= Gallons ethanol produced (wet milling or dry milling)
Flow	= Gallons wastewater generated per gallon ethanol produced
COD	= COD concentration in influent (g/l)
3.785	= Conversion factor, gallons to liters
%Plants <sub>o</sub>	= Percent of plants with onsite treatment
%WW <sub>a,p</sub>	= Percent of wastewater treated anaerobically in primary treatment
%COD <sub>p</sub>	= Percent of COD entering primary treatment
%Plants <sub>a</sub>	= Percent of plants with anaerobic secondary treatment
%Plants <sub>t</sub>	= Percent of plants with other secondary treatment
%WW <sub>a,s</sub>	= Percent of wastewater treated anaerobically in anaerobic secondary treatment
%WW <sub>a,t</sub>	= Percent of wastewater treated anaerobically in other secondary treatment
%COD <sub>s</sub>	= Percent of COD entering secondary treatment
B <sub>o</sub>	= Maximum methane producing capacity (g CH <sub>4</sub> /g COD)
MCF	= Methane correction factor
% Recovered	= Percent of wastewater treated in system with emission recovery
% Not Recovered	= 1 - percent of wastewater treated in system with emission recovery
DE	= Destruction efficiency of recovery system
1/10 <sup>9</sup>	= Conversion factor, g to kt

A time series of CH<sub>4</sub> emissions for 1990 through 2017 was developed based on production data from the Renewable Fuels Association (Cooper 2018).

*Petroleum Refining.* Petroleum refining wastewater treatment operations have the potential to produce CH<sub>4</sub> emissions from anaerobic wastewater treatment. EPA's Office of Air and Radiation performed an Information Collection Request (ICR) for petroleum refineries in 2011.<sup>8</sup> Of the responding facilities, 23.6 percent reported using non-aerated surface impoundments or other biological treatment units, both of which have the potential to lead to anaerobic conditions (ERG 2013b). In addition, the wastewater generation rate was determined to be 26.4 gallons per barrel of finished product (ERG 2013b). An average COD value in the wastewater was estimated at 0.45 kg/m<sup>3</sup> (Benyahia et al. 2006). A default MCF of 0.3 was used for partially aerobic systems.

The equation used to calculate CH<sub>4</sub> generation at petroleum refining wastewater treatment systems is presented below:

$$\text{Methane} = \text{Flow} \times \text{COD} \times \% \text{TA} \times B_o \times \text{MCF}$$

where,

Flow	= Annual flow treated through anaerobic treatment system (m <sup>3</sup> /year)
COD	= COD loading in wastewater entering anaerobic treatment system (kg/m <sup>3</sup> )

<sup>8</sup> Available online at <<https://www.epa.gov/stationary-sources-air-pollution/comprehensive-data-collected-petroleum-refining-sector>>

- 1           %TA           = Percent of wastewater treated anaerobically on site
- 2           B<sub>o</sub>           = Maximum methane producing potential of industrial wastewater (kg CH<sub>4</sub>/kg COD)
- 3           MCF           = Methane correction factor

4 A time series of CH<sub>4</sub> emissions for 1990 through 2017 was developed based on production data from the EIA 2018.  
 5 *Breweries*. Since 2010, the number of breweries has increased from less than 2,000 to greater than 6,000 (Brewers  
 6 Association 2018). This increase has primarily been driven by craft breweries, which have increased by over 250  
 7 percent during that period. Craft breweries were defined as breweries producing less than six million barrels of beer  
 8 per year, and non-craft breweries produce greater than six million barrels. With their large amount of water use and  
 9 high strength wastewater, breweries generate considerable CH<sub>4</sub> emissions from anaerobic wastewater treatment.  
 10 However, because many breweries recover their CH<sub>4</sub>, their emissions are much lower.

11 The Alcohol and Tobacco Tax and Trade Bureau (TTB) provides total beer production in barrels per year for  
 12 different facility size categories from 2007 to the present (TTB 2018). For years prior to 2007 where TTB data were  
 13 not readily available, the *Brewers Almanac* (Beer Institute 2011) was used, along with an estimated percent of craft  
 14 and non-craft breweries based on the breakdown of craft and non-craft for the years 2007 through 2017.

15 The amount of water usage by craft breweries was estimated using the Brewers Association’s *2015 Sustainability*  
 16 *Benchmarking Report* (Brewers Association 2016a) and the *2016 Benchmarking Update* (Brewers Association  
 17 2017; ERG 2018b). Non-craft brewery water usage values were from the Beverage Industry Environmental  
 18 Roundtable (BIER) benchmarking study (BIER 2017).

19 To determine the overall amount of wastewater produced, data on water use per unit of production and a  
 20 wastewater-to-water ratio were used from the Benchmarking Report (Brewers Association 2016a) for both craft and  
 21 non-craft breweries. Since brewing is a batch process, and different operations have varying organic loads, full-  
 22 strength brewery wastewater can vary widely on a day to day basis. However, the organic content of brewery  
 23 wastewater does not substantially change between craft and non-craft breweries. On average, full-strength  
 24 wastewater is about 10,600 mg/L BOD, with a typical BOD:COD ratio of 0.6 (Brewers Association 2016b). Some  
 25 breweries may collect and discharge high-strength wastewater from particular brewing processes (known as “side  
 26 streaming”) to a POTW, greatly reducing the organics content of the wastewater that is treated on site.  
 27 Subsequently, the MCF for discharge to a POTW was assumed to be zero (ERG 2018b).

28 Breweries may treat some or all of their wastewater on site prior to discharge to a POTW or receiving water. On-site  
 29 treatment operations can include physical treatment (e.g., screening, settling) which are not expected to contribute to  
 30 CH<sub>4</sub> emissions, or biological treatment, which may include aerobic treatment or pretreatment in anaerobic reactors  
 31 (ERG 2018b). The IPCC default B<sub>o</sub> of 0.25 kg CH<sub>4</sub>/kg COD and default MCFs of 0.8 for anaerobic treatment and 0  
 32 for aerobic treatment were used to estimate the CH<sub>4</sub> produced from these on-site treatment systems (IPCC 2006).  
 33 The amount of CH<sub>4</sub> recovered through anaerobic wastewater treatment was estimated, and a 99 percent destruction  
 34 efficiency was used (ERG 2018b; Stier J. 2018). Very limited activity data are available on the number of U.S.  
 35 breweries that are performing side streaming or pretreatment of wastewater prior to discharge.

36 The assumed distribution of wastewater treatment for craft and non-craft breweries are shown in Table 7-15.

37 **Table 7-15: Wastewater Treatment Distribution for Breweries**

Treatment Type	Operation Type	
	Non-Craft	Craft
Discharge to POTW with no pretreatment	0%	99%
Discharge to POTW following side streaming	0%	0.5%
Pretreatment with aerobic biological treatment	1%	0%
Pretreatment with anaerobic reactor	99%	0.5%

Source: Stier, J. (2018)

38 Methane emissions were then estimated for non-craft breweries and for craft breweries as follows:

39  
 40 
$$\text{Methane} = [(\text{Production} \times \text{Water Usage} \times \text{WW:W} \times 31)/264.172] \times \text{COD} \times ([\% \text{Plants}_{\text{potw}} \times \text{MCF}_{\text{potw}}] +$$
  
 41 
$$[\% \text{Plants}_{\text{ss}} \times \text{MCF}_{\text{potw}}] + [\% \text{Plants}_{\text{aer}} \times \text{MCF}_{\text{aer}}] + [\% \text{Plants}_{\text{a}} \times \text{MCF}_{\text{a}}]) \times \text{B}_o \times \% \text{ Not Recovered}] +$$

$$[(\text{Production} \times \text{Water Usage} \times \text{WW:W} \times 31)/264.172] \times \text{COD} \times ([\% \text{Plants}_{\text{potw}} \times \text{MCF}_{\text{potw}}] + [\% \text{Plants}_{\text{ss}} \times \text{MCF}_{\text{potw}}] + [\% \text{Plants}_{\text{aer}} \times \text{MCF}_{\text{aer}}] + [\% \text{Plants}_{\text{a}} \times \text{MCF}_{\text{a}}]) \times B_o \times (\% \text{ Recovered}) \times (1 - \text{DE}) \times 1/10^6$$

where,

4	Production	= Barrels beer produced (non-craft breweries or craft breweries)
5	Water Usage	= Barrels water utilized per barrels beer produced
6	WW:W	= Ratio, barrels of wastewater generated per barrels of water utilized
7	COD	= COD concentration in influent (kg/m <sup>3</sup> )
8	31	= Conversion factor, gallons to barrels beer
9	264.172	= Conversion factor, gallons to m <sup>3</sup>
10	%Plants <sub>potw</sub>	= Percent of plants that discharge to POTW without pretreatment
11	MCF <sub>potw</sub>	= Methane correction factor, discharge to POTW
12	%Plants <sub>ss</sub>	= Percent of plants with sidestreaming prior to POTW discharge
13	%Plants <sub>aer</sub>	= Percent of plants with primary aerobic treatment
14	MCF <sub>aer</sub>	= Methane correction factor, aerobic systems
15	%Plants <sub>a</sub>	= Percent of plants with anaerobic treatment
16	MCF <sub>a</sub>	= Methane correction factor, anaerobic systems
17	B <sub>o</sub>	= Maximum methane producing capacity (g CH <sub>4</sub> /g COD)
18	% Recovered	= Percent of wastewater treated in system with emission recovery
19	% Not Recovered	= 1 - percent of wastewater treated in system with emission recovery
20	DE	= Destruction efficiency of recovery system
21	1/10 <sup>6</sup>	= Conversion factor, kg to Gg

## 22 Domestic Wastewater N<sub>2</sub>O Emission Estimates

23 Nitrous oxide emissions from domestic wastewater (wastewater treatment) were estimated using the IPCC (2006)  
 24 methodology and supplemented with IPCC (2014) methodology to include constructed wetland emissions, including  
 25 calculations that take into account N removal with biosolids, non-consumption and industrial/commercial  
 26 wastewater N, and emissions from advanced and constructed wetlands at centralized wastewater treatment plants:

27 In the United States, a certain amount of N is removed with biosolids, which is applied to land, incinerated, or  
 28 landfilled (N<sub>SLUDGE</sub>). The value for N discharged into aquatic environments as effluent is reduced to account for the  
 29 biosolids application.

30 The *2006 IPCC Guidelines* use annual, per capita protein consumption (kg protein/person-year). For this Inventory,  
 31 the amount of protein available to be consumed is estimated based on per capita annual food availability data and its  
 32 protein content. That data are then adjusted using a factor to account for the fraction of protein actually consumed.

33 Small amounts of gaseous nitrogen oxides are formed as byproducts in the conversion of nitrate to N gas in anoxic  
 34 biological treatment systems. Approximately 7 g N<sub>2</sub>O is generated per capita per year if wastewater treatment  
 35 includes intentional nitrification and denitrification (Scheehle and Doorn 2001). Analysis of the use of treatment  
 36 systems in the United States that include denitrification has shown a significant increase in the time period between  
 37 2004 and 2012, from serving populations totaling 2.4 million people to 21.3 million people (EPA 2004 and EPA  
 38 2012). This is consistent with efforts throughout the United States to improve nutrient removal at centralized  
 39 treatment systems in response to specific water quality concerns. Based on an emission factor of 7 g per capita per  
 40 year, and data from CWNS 2004, 2008, and 2012, approximately 21.2 metric tons of additional N<sub>2</sub>O may have been  
 41 emitted via denitrification in 2004, while about 186 metric tons may have been emitted via denitrification in both  
 42 2008 and 2012. Similar analyses were completed for each year in the Inventory using data from CWNS on the  
 43 amount of wastewater in centralized systems treated in denitrification units. Plants without intentional nitrification  
 44 or denitrification are assumed to generate 3.2 g N<sub>2</sub>O per capita per year.

45 Constructed wetlands may be used as the sole treatment unit at a centralized wastewater treatment plant or may  
 46 serve as tertiary treatment after simple settling and biological treatment. Emissions from all constructed wetland  
 47 systems were included in the estimates of emissions from centralized wastewater treatment plant processes and  
 48 effluent from these plants. The emission factor of 0.0013 kg N<sub>2</sub>O-N/kg N produced for constructed wetlands is from  
 49 IPCC (2014).

1 N<sub>2</sub>O emissions from wastewater treatment plants are estimated, and as such, the N associated with these emissions is  
 2 subtracted from the amount of N estimated to be discharged into aquatic environments as effluent, consistent with  
 3 the 2006 IPCC Guidelines.

4 Nitrous oxide emissions from domestic wastewater were estimated using the following methodology:

$$5 \quad N_2O_{TOTAL} = N_2O_{PLANT} + N_2O_{EFFLUENT}$$

$$6 \quad N_2O_{PLANT} = N_2O_{NIT/DENIT} + N_2O_{WOUT\ NIT/DENIT} + N_2O_{CW\ ONLY} + N_2O_{CW\ TERTIARY}$$

$$7 \quad N_2O_{NIT/DENIT} = [(US_{POPND}) \times EF_2 \times F_{IND-COM}] \times 1/10^9$$

$$8 \quad N_2O_{WOUT\ NIT/DENIT} = \{[(US_{POP} \times WWTP) - US_{POPND} - US_{POP\ CW}] \times 10^6 \times F_{IND-COM} \times EF_1\} \times 1/10^9$$

$$9 \quad N_2O_{CW\ ONLY} = \{[(US_{POP\ CW} \times 10^6 \times Protein \times F_{NPR} \times F_{NON-CON} \times F_{IND-COM}) \times EF_4] \times 44/28\} \times 1/10^6$$

$$10 \quad N_2O_{CW\ TERTIARY} = \{[(N_{CW,INF} \times POTW\_flow\_CW \times 3.79 \times 365.25) \times EF_4] \times 44/28\} \times 1/10^6$$

$$11 \quad N_2O_{EFFLUENT} = [(US_{POP} \times WWTP \times Protein \times F_{NPR} \times F_{NON-CON} \times F_{IND-COM}) - N_{SLUDGE} - (N_2O_{PLANT} \times 10^6 \times 28/44)] \times$$

$$12 \quad EF_3 \times 44/28 \times 1/10^6$$

13 where,

14	N <sub>2</sub> O <sub>TOTAL</sub>	= Annual emissions of N <sub>2</sub> O (kt)
15	N <sub>2</sub> O <sub>PLANT</sub>	= N <sub>2</sub> O emissions from centralized wastewater treatment plants (kt)
16	N <sub>2</sub> O <sub>NIT/DENIT</sub>	= N <sub>2</sub> O emissions from centralized wastewater treatment plants with
17		nitrification/denitrification (kt)
18	N <sub>2</sub> O <sub>WOUT NIT/DENIT</sub>	= N <sub>2</sub> O emissions from centralized wastewater treatment plants without
19		nitrification/denitrification (kt)
20	N <sub>2</sub> O <sub>CW ONLY</sub>	= N <sub>2</sub> O emissions from centralized wastewater treatment plants with constructed
21		wetlands only (kt)
22	N <sub>2</sub> O <sub>CW TERTIARY</sub>	= N <sub>2</sub> O emissions from centralized wastewater treatment plants with constructed
23		wetlands used as tertiary treatment (kt)
24	N <sub>2</sub> O <sub>EFFLUENT</sub>	= N <sub>2</sub> O emissions from wastewater effluent discharged to aquatic environments (kt)
25	US <sub>POP</sub>	= U.S. population
26	US <sub>POPND</sub>	= U.S. population that is served by biological denitrification
27	US <sub>POP\ CW</sub>	= U.S. population that is served by only constructed wetland systems
28	WWTP	= Fraction of population using WWTP (as opposed to septic systems)
29	POTW_flow_CW	= Wastewater flow to POTWs that use constructed wetlands as tertiary treatment
30		(MGD)
31	EF <sub>1</sub>	= Emission factor – plants without intentional denitrification
32	EF <sub>2</sub>	= Emission factor – plant with intentional nitrification or denitrification
33	Protein	= Annual per capita protein consumption (kg/person/year)
34	N <sub>CW,INF</sub>	= Influent nitrogen concentration to constructed wetlands used as tertiary treatment
35		(mg/L)
36	F <sub>NPR</sub>	= Fraction of N in protein (kg N/kg protein)
37	F <sub>NON-CON</sub>	= Factor for non-consumed protein added to wastewater
38	F <sub>IND-COM</sub>	= Factor for industrial and commercial co-discharged protein into the sewer
39	N <sub>SLUDGE</sub>	= N removed with sludge, kg N/year
40	EF <sub>3</sub>	= Emission factor (kg N <sub>2</sub> O -N/kg sewage-N produced) – from effluent
41	EF <sub>4</sub>	= Emission factor (kg N <sub>2</sub> O -N/kg N produced) – constructed wetlands
42	3.79	= Conversion factor, gallons to liters
43	44/28	= Molecular weight ratio of N <sub>2</sub> O to N <sub>2</sub>
44	28/44	= Molecular weight ratio of N <sub>2</sub> to N <sub>2</sub> O
45	1/10 <sup>6</sup>	= Conversion factor, kg to Gg
46	1/10 <sup>9</sup>	= Conversion factor, g to Gg

47 U.S. population data were taken from the U.S. Census Bureau International Database (U.S. Census Bureau 2018)  
 48 and include the populations of the United States, American Samoa, Guam, Northern Mariana Islands, Puerto Rico,  
 49 and the U.S. Virgin Islands. The fraction of the U.S. population using wastewater treatment plants is based on data  
 50 from the 1989, 1991, 1993, 1995, 1997, 1999, 2001, 2003, 2005, 2007, 2009, 2011, 2013, and 2015 *American*

1 *Housing Survey* (U.S. Census Bureau 2015). Data for intervening years were obtained by linear interpolation and  
 2 2017 was forecasted using 1990 to 2016 data. The emission factor (EF<sub>1</sub>) used to estimate emissions from wastewater  
 3 treatment for plants without intentional nitrification or denitrification was taken from IPCC (2006), while the  
 4 emission factor (EF<sub>2</sub>) used to estimate emissions from wastewater treatment for plants with intentional nitrification  
 5 or denitrification was taken from Scheehle and Doorn (2001). The emission factor (EF<sub>4</sub>) used to estimate emissions  
 6 from surface flow constructed wetlands (0.0013 kg N<sub>2</sub>O -N/kg N produced) was taken from IPCC (2014). Data on  
 7 annual per capita protein intake were provided by the U.S. Department of Agriculture Economic Research Service  
 8 (USDA 2018b) and FAO (2018c). Protein consumption data was used directly from USDA for 1990 to 2010 and  
 9 2011 through 2013 was calculated using FAO data and a scaling factor. 2014 through 2017 were forecasted from  
 10 data for 1990 through 2013. An emission factor to estimate emissions from effluent (EF<sub>3</sub>) has not been specifically  
 11 estimated for the United States, thus the default IPCC value (0.005 kg N<sub>2</sub>O-N/kg sewage-N produced) was applied  
 12 (IPCC 2006). The fraction of N in protein (0.16 kg N/kg protein) was also obtained from IPCC (2006). The factor  
 13 for non-consumed protein (1.2) and the factor for industrial and commercial co-discharged protein (1.25) were  
 14 obtained from IPCC (2006). The amount of nitrogen removed by denitrification systems was taken from EPA  
 15 (2008a), while the population served by denitrification systems was estimated from Clean Watersheds Needs Survey  
 16 (EPA 1992, 1996, 2000, 2004, 2008b, and 2012). Sludge generation was obtained from EPA (1999) for 1988, 1996,  
 17 and 1998 and from Beecher et al. (2007) for 2004. Intervening years were interpolated and estimates for 2005  
 18 through 2017 were forecasted from the rest of the time series. The influent nitrogen concentration to constructed  
 19 wetlands used as tertiary treatment (25 mg/L) was obtained from Metcalf & Eddy (2014). An estimate for the N  
 20 removed as sludge (N<sub>SLUDGE</sub>) was obtained by determining the amount of sludge disposed by incineration, by land  
 21 application (agriculture or other), through surface disposal, in landfills, or through ocean dumping (EPA 1993b;  
 22 Beecher et al. 2007; McFarland 2001; EPA 1999). In 2017, 298 kt N was removed with sludge. Table 7-16 presents  
 23 the data for U.S. population, population served by biological denitrification, population served by wastewater  
 24 treatment plants, available protein, protein consumed, and nitrogen removed with sludge.

25 **Table 7-16: U.S. Population (Millions), Population Served by Biological Denitrification**  
 26 **(Millions), Fraction of Population Served by Wastewater Treatment (percent), Available**  
 27 **Protein (kg/person-year), Protein Consumed (kg/person-year), and Nitrogen Removed with**  
 28 **Sludge (kt-N/year)**

Year	Population	Population <sub>ND</sub>	WWTP Population	Available Protein	Protein Consumed	N Removed with Sludge
1990	253	2.0	75.6	43.1	33.2	214.2
2005	300	7.1	78.8	44.9	34.7	261.1
2013	320	19.8	81.4	43.3	33.4	285.6
2014	322	20.8	80.8	44.3	34.1	288.7
2015	325	21.8	80.2	44.3	34.1	291.8
2016	327	22.8	81.4	44.3	34.1	294.8
2017	330	23.8	81.7	44.3	34.1	297.9

Sources: Population: U.S. Census Bureau (2018); Population<sub>ND</sub>: EPA (1992), EPA (1996), EPA (2000), EPA (2004), EPA (2008b), EPA (2012); WWTP Population: U.S. Census Bureau (2015); Available Protein: USDA (2018b); N Removed with sludge: Beecher et al. (2007), McFarland (2001), EPA (1999), EPA (1993c).

## 29 Uncertainty and Time-Series Consistency

30 The overall uncertainty associated with both the 2017 CH<sub>4</sub> and N<sub>2</sub>O emission estimates from wastewater treatment  
 31 and discharge was calculated using the 2006 IPCC Guidelines Approach 2 methodology (IPCC 2006). Uncertainty  
 32 associated with the parameters used to estimate CH<sub>4</sub> emissions include that of numerous input variables used to  
 33 model emissions from domestic wastewater, and wastewater from pulp and paper manufacturing, meat and poultry  
 34 processing, fruits and vegetable processing, ethanol production, petroleum refining, and breweries. Uncertainty  
 35 associated with the parameters used to estimate N<sub>2</sub>O emissions include that of biosolids disposal, total U.S.  
 36 population, average protein consumed per person, fraction of N in protein, non-consumption nitrogen factor,  
 37 emission factors per capita and per mass of sewage-N, and for the percentage of total population using centralized  
 38 wastewater treatment plants. Uncertainty associated with constructed wetlands parameters including U.S. population  
 39 served by constructed wetlands, and emission and conversion factors are from IPCC (2014), whereas uncertainty



1 associated with POTW flow to constructed wetlands and influent BOD and nitrogen concentrations were based on  
 2 expert judgment.

3 The results of this Approach 2 quantitative uncertainty analysis are summarized in Table 7-17. Methane emissions  
 4 from wastewater treatment were estimated to be between 10.2 and 17.3 MMT CO<sub>2</sub> Eq. at the 95 percent confidence  
 5 level (or in 19 out of 20 Monte Carlo Stochastic Simulations). This indicates a range of approximately 28 percent  
 6 below to 21 percent above the 2017 emissions estimate of 14.3 MMT CO<sub>2</sub> Eq. Nitrous oxide emissions from  
 7 wastewater treatment were estimated to be between 1.2 and 10.3 MMT CO<sub>2</sub> Eq., which indicates a range of  
 8 approximately 75 percent below to 108 percent above the 2017 emissions estimate of 5.0 MMT CO<sub>2</sub> Eq.

9 **Table 7-17: Approach 2 Quantitative Uncertainty Estimates for CH<sub>4</sub> Emissions from**  
 10 **Wastewater Treatment (MMT CO<sub>2</sub> Eq. and Percent)**

Source	Gas	2017 Emission Estimate (MMT CO <sub>2</sub> Eq.)	Uncertainty Range Relative to Emission Estimate <sup>a</sup> (%)			
			Lower Bound	Upper Bound	Lower Bound	Upper Bound
<b>Wastewater Treatment</b>	<b>CH<sub>4</sub></b>	<b>14.3</b>	<b>10.2</b>	<b>17.3</b>	<b>-28%</b>	<b>+21%</b>
Domestic	CH <sub>4</sub>	8.6	6.1	10.3	-29%	+21%
Industrial	CH <sub>4</sub>	5.7	2.9	8.7	-49%	+51%
<b>Wastewater Treatment</b>	<b>N<sub>2</sub>O</b>	<b>5.0</b>	<b>1.2</b>	<b>10.3</b>	<b>-75%</b>	<b>+108%</b>

<sup>a</sup> Range of emission estimates predicted by Monte Carlo Stochastic Simulation for a 95 percent confidence interval.

## 11 QA/QC and Verification

12 General QA/QC procedures were applied to activity data, documentation, and emission calculations consistent with  
 13 the U.S. QA/QC plan, which is in accordance with Vol. 1 Chapter 6 of *2006 IPCC Guidelines* (see Annex 8 for  
 14 more details). This effort included a general or Tier 1 analysis, including the following checks:

- 15 • Checked for transcription errors in data input;
- 16 • Ensured references were specified for all activity data used in the calculations;
- 17 • Checked a sample of each emission calculation used for the source category;
- 18 • Checked that parameter and emission units were correctly recorded and that appropriate conversion factors  
 19 were used;
- 20 • Checked for temporal consistency in time series input data for each portion of the source category;
- 21 • Confirmed that estimates were calculated and reported for all portions of the source category and for all  
 22 years;
- 23 • Investigated data gaps that affected trends of emissions estimates; and
- 24 • Compared estimates to previous estimates to identify significant changes.

25 All transcription errors identified were corrected and documented. The QA/QC analysis did not reveal any systemic  
 26 inaccuracies or incorrect input values.

## 27 Recalculations Discussion

28 Population data were updated to reflect revised U.S. Census Bureau datasets which resulted in changes to 1991  
 29 through 2016 values (U.S. Census Bureau 2018). Forecasted protein data was updated which resulted in changes to  
 30 2014 through 2016 available protein and protein consumed values.

31 EPA evaluated pulp and paper production, average BOD concentrations in raw wastewater, and COD:BOD ratio  
 32 based on the National Council of Air and Stream Improvement's (NCASI) recommendation and determined updates  
 33 to current Inventory data were appropriate. EPA updated production values from summing wood pulp and paper and  
 34 paperboard to summing market pulp and paper and paperboard production which resulted in changes for the entire  
 35 time series. This change also resulted in an update to the data source for pulp and paper production prior to 2002  
 36 from the Lockwood-Post to FAO. EPA updated raw wastewater BOD concentrations and the COD:BOD ratio of

1 influent wastewater based on industry data provided by NCASI (Malmberg 2018) which resulted in changes for  
2 1999 through 2016 and the entire time series, respectively.

3 EPA evaluated domestic raw BOD production and determined updates to current Inventory data were appropriate to  
4 reflect differences in waste characteristics from households with and without kitchen garbage disposals. The BOD<sub>5</sub>  
5 production rate was determined using BOD generation rates per capita both with and without kitchen scraps  
6 (Metcalf & Eddy 2003; Metcalf & Eddy 2014) as well as an estimated percent of housing units that utilize kitchen  
7 garbage disposals (ERG 2018a). In addition to applying the distinction of with and without kitchen scraps between  
8 BOD generation rates per capita, the value for the BOD generation rate changed with an updated source (Metcalf &  
9 Eddy 2014). This update further impacted the amount of domestic BOD produced from 2004 through 2016. EPA  
10 now estimates a dynamic BOD generation rate per capita which resulted in changes for the entire times series (ERG  
11 2018a).

12 On an ongoing basis, EPA reviews other industries that have the potential to emit CH<sub>4</sub> from their wastewater  
13 treatment systems because they treat wastewater with significant organics loads. EPA evaluated emissions estimates  
14 from wastewater treatment processes at breweries for potential inclusion in the Inventory. Based on data from the  
15 Brewers Association (Brewers Association 2018; Brewers Association 2017; Brewers Association 2016a; Brewers  
16 Association 2016b), the Beverage Industry Environmental Roundtable (BIER 2017), the Alcohol and Tobacco Tax  
17 and Trade Bureau (TTB 2018), and conversations with industry experts as described above, EPA determined that  
18 this industry generates significant quantities of CH<sub>4</sub> from wastewater treatment operations, though a majority of the  
19 emissions are recovered. As a result, EPA determined that the brewery industry is an appropriate category to include  
20 in the Inventory.

21 EPA was also made aware of an error in the emissions calculation for centrally treated aerobic systems using  
22 constructed wetlands as tertiary treatment after this draft was compiled for public review, so this error will be  
23 corrected following public review. This correction will affect the entire time series and is expected to result in an  
24 average decrease of 0.5% for domestic methane emissions and 0.3% for total methane emissions across the time  
25 series.

## 26 **Planned Improvements**

27 EPA will continue to investigate the following improvements to the wastewater emissions estimates in the  
28 Inventory:

- 29 • Continue working with the NCASI to further refine the market pulp production values as well as update  
30 wastewater characteristic data as new or improved data become available;
- 31 • Investigate updated sources of activity data for wastewater treatment system type to distinguish between  
32 aerobic, anaerobic, and other systems with the potential to generate CH<sub>4</sub>. This includes re-evaluating a  
33 methodology that was developed so that the 2008 and 2012 CWNS data could be used in estimating  
34 emissions from constructed wetlands to determine if it could be extended to all types of systems; and
- 35 • Continue reviewing other industrial wastewater treatment sources for those industries believed to discharge  
36 significant loads of BOD or COD, including dairy products processing.

37 *In addition, EPA will continue to monitor potential sources for updating Inventory data, including:*

- 38 • Sources of data for updating the factor for industrial and commercial co-discharged protein to determine if  
39 the IPCC factor currently used (1.25) is underestimating the contribution of industrial wastewater to N<sub>2</sub>O  
40 emissions;
- 41 • WEF biosolid data as a potential source of digester, sludge, and biogas data from POTWs;
- 42 • Reports based on international research and other countries' inventory submissions to inform potential  
43 updates to the Inventory's emission factors, methodologies, or included industries;
- 44 • Research by groups such as the Water Research Foundation (formerly Water Environment Research  
45 Federation) on emissions from various types of municipal treatment systems, country-specific N<sub>2</sub>O  
46 emission factors, and flare efficiencies and data that indicate septic soil systems are a source of N<sub>2</sub>O for the  
47 potential development of appropriate emission factors for septic system N<sub>2</sub>O emissions;
- 48 • Sources of data for development of a country-specific methodology for N<sub>2</sub>O emissions associated with on-  
49 site industrial wastewater treatment operations, including the appropriateness of using IPCC's default factor  
50 for domestic wastewater (0.005 kg N<sub>2</sub>O-N/kg N);

- Additional data sources for stand-alone centralized waste treaters. These data may inform current treatment assumptions for industrial categories;
- Additional data sources for improving the uncertainty of the estimate of N entering municipal treatment systems; and
- Data to update the value used for N content of sludge, the amount of sludge produced, and sludge disposal practices, along with increasing the transparency of the fate of sludge produced in wastewater treatment.

A refinement of the *2006 IPCC Guidelines* is currently underway to incorporate abundant new scientific and empirical knowledge published since 2006 which the IPCC should take into account, particularly with respect to data for emission factor development. For wastewater treatment, this refinement includes a review of methane and nitrous oxide emission factors, and an assessment of adding methodologies to account for nitrous oxide emissions from both domestic and industrial wastewater. EPA will continue to monitor the status of this refinement for potential updates to the wastewater inventory methodology.

These planned improvements were described in greater detail in the previous Inventory report; please see Section 7.2 of the *Inventory of U.S. Greenhouse Gas Emissions and Sinks: 1990-2016*.

## 7.3 Composting (CRF Source Category 5B1)

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Composting of organic waste, such as food waste, garden (yard) and park waste, and wastewater treatment sludge and/or biosolids, is common in the United States. Advantages of composting include reduced volume of the waste, stabilization of the waste, and destruction of pathogens in the waste. The end products of composting, depending on its quality, can be recycled as a fertilizer and soil amendment, or be disposed of in a landfill.

Composting is an aerobic process and a large fraction of the degradable organic carbon in the waste material is converted into carbon dioxide (CO<sub>2</sub>). Methane (CH<sub>4</sub>) is formed in anaerobic sections of the compost, which are created when there is excessive moisture or inadequate aeration (or mixing) of the compost pile. This CH<sub>4</sub> is then oxidized to a large extent in the aerobic sections of the compost. The estimated CH<sub>4</sub> released into the atmosphere ranges from less than 1 percent to a few percent of the initial C content in the material (IPCC 2006). Depending on how well the compost pile is managed, nitrous oxide (N<sub>2</sub>O) emissions can be produced. The formation of N<sub>2</sub>O depends on the initial nitrogen content of the material and is mostly due to nitrogen oxide (NO<sub>x</sub>) denitrification during the thermophilic and secondary mesophilic stages of composting (Cornell 2007). Emissions vary and range from less than 0.5 percent to 5 percent of the initial nitrogen content of the material (IPCC 2006). Animal manures are typically expected to generate more N<sub>2</sub>O than, for example, yard waste, however data are limited.

From 1990 to 2017, the amount of waste composted in the United States increased from 3,810 kt to 21,333 kt. There was some fluctuation in the amount of waste composted between 2006 to 2009. A peak of 20,049 kt composted was observed in 2008, followed by a steep drop the following year to 18,824 kt composted, presumably driven by the economic crisis. Since then, the amount of waste composted has gradually increased, and when comparing 2010 to 2017, a 16.6 percent increase in waste composted is observed. Emissions of CH<sub>4</sub> and N<sub>2</sub>O from composting from 2010 to 2017 have increased by the same percentage. In 2017, CH<sub>4</sub> emissions from composting (see Table 7-18 and Table 7-19) were 2.2 MMT CO<sub>2</sub> Eq. (86 kt), and N<sub>2</sub>O emissions from composting were 1.9 MMT CO<sub>2</sub> Eq. (6 kt), representing a slight increase compared to 2016. The wastes composted primarily include yard trimmings (grass, leaves, and tree and brush trimmings) and food scraps from the residential and commercial sectors (such as grocery stores; restaurants; and school, business, and factory cafeterias). The composted waste quantities reported here do not include backyard composting or agricultural composting.

The growth in composting since the 1990s and specifically over the past decade is attributable primarily to the following factors: (1) the enactment of legislation by state and local governments that discouraged the disposal of yard trimmings and food waste in landfills, (2) yard trimming collection and yard trimming drop off sites provided by local solid waste management districts/divisions, (3) an increased awareness of the environmental benefits of composting, and (4) loans or grant programs to establish or expand composting infrastructure. Most bans or diversion laws on the disposal of yard trimmings were initiated in the early 1990s by state or local governments (U.S. Composting Council 2010). California, for example, enacted a waste diversion law for organics including yard trimmings and food scraps in 1999 (AB939) that required jurisdictions to divert 50 percent of the waste stream by 2000, or be subjected to fines. By 2010, 25 states, representing about 50 percent of the nation's population, had

1 enacted such legislation (ILSR 2014; BioCycle 2010). There are many more initiatives at the metro and municipal  
 2 level across the United States. More than 3,280 composting facilities exist in the United States with most (71  
 3 percent) composting yard trimmings only (ISLR 2014).

4 **Table 7-18: CH<sub>4</sub> and N<sub>2</sub>O Emissions from Composting (MMT CO<sub>2</sub> Eq.)**

Activity	1990	2005	2013	2014	2015	2016	2017
CH <sub>4</sub>	0.4	1.9	2.0	2.1	2.1	2.1	2.2
N <sub>2</sub> O	0.3	1.7	1.8	1.9	1.9	1.9	1.9
<b>Total</b>	<b>0.7</b>	<b>3.5</b>	<b>3.9</b>	<b>4.0</b>	<b>4.0</b>	<b>4.0</b>	<b>4.1</b>

5 **Table 7-19: CH<sub>4</sub> and N<sub>2</sub>O Emissions from Composting (kt)**

Activity	1990	2005	2013	2014	2015	2016	2017
CH <sub>4</sub>	15	75	81	84	85	85	86
N <sub>2</sub> O	1	6	6	6	6	6	7

## 6 Methodology

7 Methane and N<sub>2</sub>O emissions from composting depend on factors such as the type of waste composted, the amount  
 8 and type of supporting material (such as wood chips and peat) used, temperature, moisture content (e.g., wet and  
 9 fluid versus dry and crumbly), and aeration during the composting process.

10 The emissions shown in Table 7-18 and Table 7-19 were estimated using the IPCC default (Tier 1) methodology  
 11 (IPCC 2006), which is the product of an emission factor and the mass of organic waste composted (note: no CH<sub>4</sub>  
 12 recovery is expected to occur at composting operations in the emission estimates presented):

13  
 14 where,

15  $E_i$  = CH<sub>4</sub> or N<sub>2</sub>O emissions from composting, kt CH<sub>4</sub> or N<sub>2</sub>O,  
 16  $M$  = mass of organic waste composted in kt,  
 17  $EF_i$  = emission factor for composting, 4 t CH<sub>4</sub>/kt of waste treated (wet basis) and  
 18 0.3 t N<sub>2</sub>O/kt of waste treated (wet basis) (IPCC 2006), and  
 19  $i$  = designates either CH<sub>4</sub> or N<sub>2</sub>O.

20 Estimates of the quantity of waste composted ( $M$ ) are presented in Table 7-20 for select years. Estimates of the  
 21 quantity composted for 1990, 2005, 2010, and 2014 to 2015 were taken from EPA's *Advancing Sustainable*  
 22 *Materials Management: Facts and Figures 2015* (EPA 2018); the estimate of the quantity composted for 2012 to  
 23 2013 was taken from EPA's *Advancing Sustainable Materials Management: Facts and Figures 2014* report; the  
 24 estimate of the quantity composted for 2011 was taken from EPA's *Municipal Solid Waste In The United States:*  
 25 *2012 Facts and Figures* (EPA 2014); estimates of the quantity composted for 2016 and 2017 were extrapolated  
 26 using the 2015 quantity composted and a ratio of the U.S. population growth between 2015 and 2016, and 2016 to  
 27 2017 (U.S. Census Bureau 2016, 2017, and 2018).

28 **Table 7-20: U.S. Waste Composted (kt)**

Activity	1990	2005	2013	2014	2015	2016	2017
Waste Composted	3,810	18,643	20,358	20,884	21,219	21,332	21,503

## 29 Uncertainty and Time-Series Consistency

30 The estimated uncertainty from the 2006 IPCC Guidelines is ±50 percent for the Tier 1 methodology. Emissions  
 31 from composting in 2017 were estimated to be between 2.0 and 6.1 MMT CO<sub>2</sub> Eq., which indicates a range of 50  
 32 percent below to 50 percent above the actual 2017 emission estimate of 4.1 MMT CO<sub>2</sub> Eq. (see Table 7-21).

$$E_i = M \times EF_i$$

**Table 7-21: Tier 1 Quantitative Uncertainty Estimates for Emissions from Composting (MMT CO<sub>2</sub> Eq. and Percent)**

Source	Gas	2017 Emission Estimate (MMT CO <sub>2</sub> Eq.)	Uncertainty Range Relative to Emission Estimate (MMT CO <sub>2</sub> Eq.)			
			Lower Bound	Upper Bound	Lower Bound (%)	Upper Bound (%)
Composting	CH <sub>4</sub> , N <sub>2</sub> O	4.1	2.0	6.1	-50%	+50%

## QA/QC and Verification

General QA/QC procedures were applied to data gathering and input, documentation, and calculations consistent with the U.S. QA/QC plan, which is in accordance with Vol. 1 Chapter 6 of *2006 IPCC Guidelines* (see Annex 8 for more details).

## Recalculations Discussion

Emissions recalculations were made in this Inventory year for the years 2015 and 2016 per the release of the EPA *Advancing Sustainable Materials Management: 2015 Facts and Figures* report (EPA 2018). The tonnage of waste composted for the year 2015 was previously extrapolated based on the tonnage composted in EPA’s *Advancing Sustainable Materials Management: 2014 Facts and Figures* report for the year 2014 and a ratio of U.S. population growth between 2014 and 2015. Because of this change to the 2015 composting tonnage, the extrapolated tonnage for the year 2016 was also altered. Table 7-19 has been updated to reflect the changes in composting emissions as a result of these updated tonnage values.

## Planned Improvements

For future Inventories, additional efforts will be made to improve the estimates of CH<sub>4</sub> and N<sub>2</sub>O emissions from composting. For example, a literature search on emission factors and composting systems and management techniques has been completed and will be documented in a technical memorandum for the 1990 through 2017 Inventory. The purpose of this literature review was to compile all published emission factors specific to various composting systems and composted materials. This information will be used to determine whether the emission factors used in the current methodology should be revised, or expanded to account for geographical differences and/or differences in composting systems used. For example, outdoor composting processes in arid regions typically require the addition of moisture compared to similar composting processes in wetter climates. Additionally, composting systems that primarily compost food waste may generate CH<sub>4</sub> at different rates than those that compost yard trimmings because the food waste may have a higher moisture content and more readily degradable material. Further investigation into accounting of composting emissions estimates across other applicable sections of the Inventory, in cooperation with the LULUCF Settlements section, will also be completed.

Additional efforts are being made to improve the comprehensiveness of the composting Inventory by incorporating composted waste from U.S. territories. EPA conducted a desk-based investigation into industrial/commercial composting facilities in the U.S. territories and identified facilities in Puerto Rico. Additional efforts are being made to collect information on the year the identified facilities began operating, an estimate of the quantity of waste composted, and approximate land area or population (or households) the facilities serve. This data may be incorporated into the current or future Inventories as a methodological improvement.

## 7.4 Waste Incineration (CRF Source Category 5C1)

As stated earlier in this chapter, carbon dioxide (CO<sub>2</sub>), nitrous oxide (N<sub>2</sub>O), and methane (CH<sub>4</sub>) emissions from the incineration of waste are accounted for in the Energy sector rather than in the Waste sector because almost all incineration of municipal solid waste (MSW) in the United States occurs at waste-to-energy facilities where useful energy is recovered. Similarly, the Energy sector also includes an estimate of emissions from burning waste tires and hazardous industrial waste, because virtually all of the combustion occurs in industrial and utility boilers that recover energy. The incineration of waste in the United States in 2017 resulted in 11.1 MMT CO<sub>2</sub> Eq. of emissions, over half of which (6.2 MMT CO<sub>2</sub> Eq.) is attributable to the combustion of plastics. For more details on emissions from the incineration of waste, see Section 3.3 of the Energy chapter.

Additional sources of emissions from waste incineration include non-hazardous industrial waste incineration and medical waste incineration. As described in Annex 5 of this report, data are not readily available for these sources and emission estimates are not provided. An analysis of the likely level of emissions was conducted based on a 2009 study of hospital/ medical/ infectious waste incinerator (HMIWI) facilities in the United States (RTI 2009). Based on that study's information of waste throughput and an analysis of the fossil-based composition of the waste, it was determined that annual greenhouse gas emissions for medical waste incineration would be below 500 kt CO<sub>2</sub> Eq. per year and considered insignificant for the purposes of Inventory reporting under the UNFCCC. More information on this analysis is provided in Annex 5.

## 7.5 Waste Sources of Precursor Greenhouse Gases

In addition to the main greenhouse gases addressed above, waste generating and handling processes are also sources of precursor gases. The reporting requirements of the UNFCCC<sup>9</sup> request that information be provided on precursor greenhouse gases, which include carbon monoxide (CO), nitrogen oxides (NO<sub>x</sub>), non-CH<sub>4</sub> volatile organic compounds (NMVOCs), and sulfur dioxide (SO<sub>2</sub>). These gases are not direct greenhouse gases, but indirectly affect terrestrial radiation absorption by influencing the formation and destruction of tropospheric and stratospheric ozone, or, in the case of SO<sub>2</sub>, by affecting the absorptive characteristics of the atmosphere. Additionally, some of these gases may react with other chemical compounds in the atmosphere to form compounds that are greenhouse gases. Total emissions of NO<sub>x</sub>, CO, and NMVOCs from waste sources for the years 1990 through 2017 are provided in Table 7-22. Sulfur dioxide emissions are presented in Section 2.3 of the Trends chapter and Annex 6.3.

**Table 7-22: Emissions of NO<sub>x</sub>, CO, and NMVOC from Waste (kt)**

Gas/Source	1990	2005	2013	2014	2015	2016	2017
<b>NO<sub>x</sub></b>	+	2	2	2	2	2	2
Landfills	+	2	2	2	2	2	2
Wastewater Treatment	+	0	0	0	0	0	0
Miscellaneous <sup>a</sup>	+	0	0	0	0	0	0
<b>CO</b>	1	7	7	8	8	8	8
Landfills	1	6	7	8	8	8	8
Wastewater Treatment	+	+	+	1	1	1	1
Miscellaneous <sup>a</sup>	+	0	0	0	0	0	0
<b>NMVOCs</b>	<b>673</b>	<b>114</b>	<b>58</b>	<b>68</b>	<b>68</b>	<b>68</b>	<b>68</b>
Wastewater Treatment	57	49	25	29	29	29	29

<sup>9</sup> See <<http://unfccc.int/resource/docs/2013/cop19/eng/10a03.pdf>>.

Miscellaneous <sup>a</sup>	557	43	22	26	26	26	26
Landfills	58	22	11	13	13	13	13

+ Does not exceed 0.5 kt.

<sup>a</sup> Miscellaneous includes TSDFs (Treatment, Storage, and Disposal Facilities under the Resource Conservation and Recovery Act [42 U.S.C. § 6924, SWDA § 3004]) and other waste categories.

Note: Totals may not sum due to independent rounding.

## 1 Methodology

2 Emission estimates for 1990 through 2017 were obtained from data published on the National Emission Inventory  
3 (NEI) Air Pollutant Emission Trends web site (EPA 2018), and disaggregated based on EPA (2003). Emission  
4 estimates of these gases were provided by sector, using a “top down” estimating procedure—emissions were  
5 calculated either for individual sources or for many sources combined, using basic activity data (e.g., the amount of  
6 raw material processed) as an indicator of emissions. National activity data were collected for individual categories  
7 from various agencies. Depending on the category, these basic activity data may include data on production, fuel  
8 deliveries, raw material processed, etc.

## 9 Uncertainty and Time-Series Consistency

10 No quantitative estimates of uncertainty were calculated for this source category. Methodological recalculations  
11 were applied to the entire time series to ensure time-series consistency from 1990 through 2017. Details on the  
12 emission trends through time are described in more detail in the Methodology section, above.