

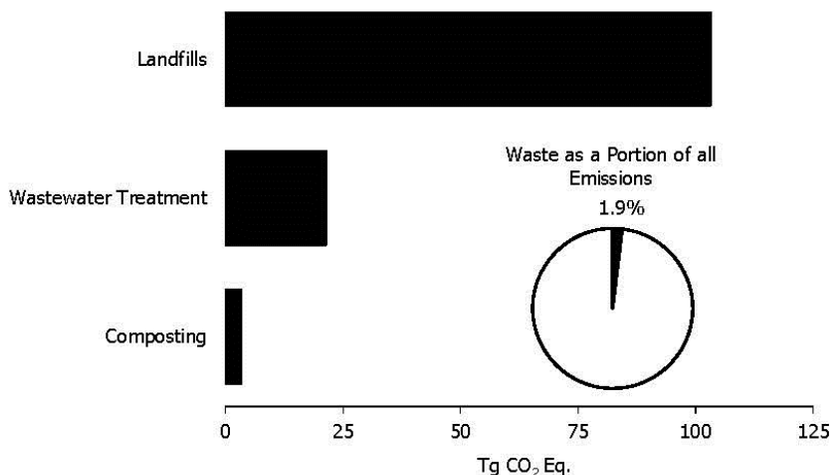
8. Waste

Waste management and treatment activities are sources of greenhouse gas emissions (see Figure 8-1). Landfills accounted for approximately 17.7 percent of total U.S. anthropogenic methane (CH₄) emissions in 2011, the second largest contribution of any CH₄ source in the United States. Additionally, wastewater treatment and composting of organic waste accounted for approximately 2.8 percent and less than 1 percent of U.S. methane emissions, respectively. Nitrous oxide (N₂O) emissions from the discharge of wastewater treatment effluents into aquatic environments were estimated, as were N₂O emissions from the treatment process itself. N₂O emissions from composting were also estimated. Together, these waste activities account for less than 2 percent of total U.S. N₂O emissions. Nitrogen oxides (NO_x), carbon monoxide (CO), and non-CH₄ 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 8-1 and Table 8-2.

CO₂, N₂O, and CH₄ 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, because virtually all of the combustion occurs in industrial and utility boilers that recover energy. The incineration of waste in the United States in 2011 resulted in 12.4 Tg CO₂ Eq. emissions, nearly half of which is attributable to the combustion of plastics. For more details on emissions from the incineration of waste, see Section 3.3.

Figure 8-1: 2010 Waste Chapter Greenhouse Gas Sources



[BEGIN BOX]

Box 8-1: Methodological approach for estimating and reporting U.S. emissions and sinks

In following the UNFCCC requirement under Article 4.1 to develop and submit national greenhouse gas emission inventories, the emissions and sinks presented in this report and this chapter, are organized by source and sink categories and calculated using internationally-accepted methods provided by the Intergovernmental Panel on Climate Change (IPCC).²⁴³ Additionally, the calculated emissions and sinks in a given year for the United States are presented in a common manner in line with the UNFCCC reporting guidelines for the reporting of inventories

²⁴³ See <http://www.ipcc-nggip.iges.or.jp/public/index.html>.

1 under this international agreement.²⁴⁴ The use of consistent methods to calculate emissions and sinks by all nations
 2 providing their inventories to the UNFCCC ensures that these reports are comparable. In this regard, U.S. emissions
 3 and sinks reported in this inventory report are comparable to emissions and sinks reported by other countries.
 4 Emissions and sinks provided in this inventory do not preclude alternative examinations,²⁴⁵ but rather this inventory
 5 presents emissions and sinks in a common format consistent with how countries are to report inventories under the
 6 UNFCCC. The report itself, and this chapter, follows this standardized format, and provides an explanation of the
 7 IPCC methods used to calculate emissions and sinks, and the manner in which those calculations are conducted.

8 [END BOX]

10 Overall, in 2011, waste activities generated emissions of 127.6 Tg CO₂ Eq., or just under 2 percent of total U.S.
 11 greenhouse gas emissions.

13 Table 8-1: Emissions from Waste (Tg CO₂ Eq.)

Gas/Source	1990	2005	2007	2008	2009	2010	2011
CH₄	164.0	130.6	129.9	131.9	131.4	124.7	120.7
Landfills	147.8	112.5	111.6	113.6	113.3	106.8	103.0
Wastewater Treatment	15.9	16.5	16.6	16.6	16.5	16.4	16.2
Composting	0.3	1.6	1.7	1.7	1.6	1.5	1.5
N₂O	3.8	6.4	6.7	6.8	6.7	6.8	6.9
Domestic Wastewater Treatment	3.5	4.7	4.8	4.9	5.0	5.1	5.2
Composting	0.4	1.7	1.8	1.9	1.8	1.7	1.7
Total	167.8	136.9	136.5	138.7	138.1	131.4	127.6

Note: Totals may not sum due to independent rounding.

14 Table 8-2: Emissions from Waste (Gg)

Gas/Source	1990	2005	2007	2008	2009	2010	2011
CH₄	7,810	6,217	6,184	6,281	6,258	5,936	5,750
Landfills	7,037	5,357	5,314	5,409	5,397	5,084	4,906
Wastewater Treatment	758	785	791	791	786	779	770
Composting	15	75	79	80	75	73	74
N₂O	12	21	21	22	22	22	22
Domestic Wastewater Treatment	11	15	16	16	16	16	17
Composting	1	6	6	6	6	5	6

Note: Totals may not sum due to independent rounding.

15 [BEGIN BOX]

16 Box 8-2: Waste Data from the Greenhouse Gas Reporting Program – TO BE UPDATED

On October 30, 2009, the U.S. EPA published a rule for the mandatory reporting of greenhouse gases from large GHG emissions sources in the United States. Implementation of 40 CFR Part 98 is referred to as EPA's Greenhouse Gas Reporting Program (GHGRP). 40 CFR part 98 applies to direct greenhouse gas emitters, fossil fuel suppliers,

²⁴⁴ See http://unfccc.int/national_reports/annex_i_ghg_inventories/national_inventories_submissions/items/5270.php.

²⁴⁵ For example, see <http://www.epa.gov/aboutepa/oswer.html>.

industrial gas suppliers, and facilities that inject CO₂ underground for sequestration or other reasons and requires reporting by 41 industrial categories. Reporting is at the facility level, except for certain suppliers of fossil fuels and industrial greenhouse gases. In general, the threshold for reporting is 25,000 metric tons or more of CO₂ Eq. per year. For calendar year 2010, the first year in which data were reported, facilities in 29 categories provided in 40 CFR part 98 were required to report their 2010 emissions by the September 30, 2011 reporting deadline.

EPA's GHGRP dataset and the data presented in this inventory report are complementary and, as indicated in the respective planned improvements sections for source categories in this chapter, EPA is analyzing how to use facility-level GHGRP data to improve the national estimates presented in this inventory. Most methodologies used in EPA's GHGRP are consistent with IPCC, though for EPA's GHGRP, facilities collect detailed information specific to their operations according to detailed measurement standards. This may differ with the more aggregated data collected for the inventory to estimate total, national U.S. emissions. In addition, it should be noted that the definitions and provisions for reporting fuel types in EPA's GHGRP may differ from those used in the national inventory in meeting the UNFCCC reporting guidelines. In line with the UNFCCC reporting guidelines²⁴⁶, the inventory report is a comprehensive accounting of all emissions from fuel types identified in the IPCC guidelines and provides a separate reporting of emissions from biomass. Further information on the reporting categorizations in EPA's GHGRP and specific data caveats associated with monitoring methods in EPA's GHGRP has been provided on the EPA's GHGRP website.²⁴⁷

EPA presents the data collected by EPA's GHGRP through a data publication tool²⁴⁸ that allows data to be viewed in several formats including maps, tables, charts and graphs for individual facilities or groups of facilities.

1 [END BOX]

2 **8.1. Landfills (IPCC Source Category 6A1)**

3 In the United States, solid waste is managed by landfilling, recovery through recycling or composting, and
4 combustion through waste-to-energy facilities. Disposing of solid waste in modern, managed landfills is the most
5 commonly used waste management technique in the United States. More information on how solid waste data are
6 collected and managed in the United States is provided in Box 8-3 and Box 8-4. The municipal solid waste (MSW)
7 and industrial waste landfills referred to in this section are all modern landfills that must comply with a variety of
8 regulations as discussed in Box 8-5. Disposing of waste in illegal dumping sites is not considered to have occurred
9 in years later than 1980 and these sites are not considered to contribute to net emissions in this section for the
10 inventory time frame of 1990 to 2011. MSW landfills, or sanitary landfills, are sites where MSW is managed to
11 prevent or minimize health, safety, and environmental impacts. Waste is deposited in different cells and covered
12 daily with soil; many have environmental monitoring systems to track performance, collect leachate, and collect
13 landfill gas. Industrial waste landfills are constructed in a similar way as MSW landfills, but accept waste produced
14 by industrial activity, such as factories, mills, and mines.

15 After being placed in a landfill, organic waste (such as paper, food scraps, and yard trimmings) is initially
16 decomposed by aerobic bacteria. After the oxygen has been depleted, the remaining waste is available for
17 consumption by anaerobic bacteria, which break down organic matter into substances such as cellulose, amino acids,
18 and sugars. These substances are further broken down through fermentation into gases and short-chain organic
19 compounds that form the substrates for the growth of methanogenic bacteria. These methane- (CH₄) producing
20 anaerobic bacteria convert the fermentation products into stabilized organic materials and biogas consisting of
21 approximately 50 percent biogenic carbon dioxide (CO₂) and 50 percent CH₄, by volume. Landfill biogas also
22 contains trace amounts of non-methane organic compounds (NMOC) and volatile organic compounds (VOC) that
23 either result from decomposition by-products or volatilization of biodegradable wastes (EPA 2008).

24 Methane and CO₂ are the primary constituents of landfill gas generation and emissions. However, the 2006
25 Intergovernmental Panel on Climate Change (IPCC) Guidelines set an international convention to not report

²⁴⁶ See <http://unfccc.int/resource/docs/2006/sbsta/eng/09.pdf>.

²⁴⁷ See

<<http://www.ccdsupport.com/confluence/display/ghgp/Detailed+Description+of+Data+for+Certain+Sources+and+Processes>>.

²⁴⁸ See <<http://ghgdata.epa.gov>>.

1 biogenic CO₂ released due to landfill decomposition in the Waste sector (IPCC 2006). Carbon dioxide emissions
2 are estimated and reported for under the Land Use/Land Use Change and Forestry (LULUCF) sector (see Box 8-6).
3 Additionally, emissions of NMOC and VOC are not estimated because they are considered to be emitted in trace
4 amounts. Nitrous oxide (N₂O) emissions from the disposal and application of sewage sludge on landfills are also not
5 explicitly modeled as part of greenhouse gas emissions from landfills. N₂O emissions from sewage sludge applied
6 to landfills as a daily cover or for disposal are expected to be relatively small because the microbial environment in
7 an anaerobic landfill is not very conducive to the nitrification and denitrification processes that result in N₂O
8 emissions. Furthermore, the 2006 IPCC Guidelines (IPCC 2006) did not include a methodology for estimating N₂O
9 emissions from solid waste disposal sites “because they are not significant.” Therefore, only CH₄ generation and
10 emissions are estimated for landfills under the Waste sector.

11 Methane generation and emissions from landfills are a function of several factors, including: (1) the total amount of
12 waste-in-place, which is the total waste landfilled annually over the operational lifetime of a landfill; (2) the
13 characteristics of the landfill receiving waste (e.g., composition of waste-in-place, size, climate, cover material); (3)
14 the amount of CH₄ that is recovered and either flared or used for energy purposes; and (4) the amount of CH₄
15 oxidized as the landfill gas passes through the cover material into the atmosphere. Each landfill has unique
16 characteristics, but all managed landfills practice similar operating practices, including the application of a daily and
17 intermediate cover material over the waste being disposed of in the landfill to prevent odor and reduce risks to
18 public health. Based on recent literature, the specific type of cover material used can affect the rate of oxidation of
19 landfill gas (RTI 2011). The most commonly used cover materials are soil, clay, and sand. Some states also permit
20 the use of green waste, tarps, waste derived materials, sewage sludge or biosolids, and contaminated soil as a daily
21 cover. Methane production typically begins one or two years after waste is disposed of in a landfill and will continue
22 for 10 to 60 years or longer as the degradable waste decomposes over time.

23 In 2011, landfill CH₄ emissions were approximately 103.0 Tg CO₂ Eq. (4,906 Gg of CH₄), representing the third
24 largest source of CH₄ emissions in the United States, behind natural gas systems and enteric fermentation.
25 Emissions from MSW landfills, which received about 69 percent of the total solid waste generated in the United
26 States, accounted for about 94 percent of total landfill emissions, while industrial landfills accounted for the
27 remainder. Approximately 1,900 to 2,000 operational MSW landfills exist in the United States, with the largest
28 landfills receiving most of the waste and generating the majority of the CH₄ emitted (EPA 2010; *BioCycle* 2010;
29 WBJ 2010). Conversely, there are approximately 3,200 MSW landfills in the United States that have been closed
30 since 1980 (for which a closure data is known, WBJ 2010). While the number of active MSW landfills has
31 decreased significantly over the past 20 years, from approximately 6,326 in 1990 to approximately 2,000 in 2010,
32 the average landfill size has increased (EPA 2010; *BioCycle* 2010; WBJ 2010). The exact number of active and
33 closed dedicated industrial waste landfills is not known at this time, but the Waste Business Journal total of landfills
34 that accept industrial and construction and demolition debris for 2010 is 1,305 (WBJ, 2010).

35 The estimated annual quantity of waste placed in MSW landfills increased 26 percent from about 205 Tg in 1990 to
36 258 Tg in 2011 (see Annex 3.13). Net CH₄ emissions have fluctuated from year to year, but a slowly decreasing
37 trend has been observed over the past decade despite increased waste disposal amounts. For example, from 1990 to
38 2011, net CH₄ emissions from landfills decreased by approximately 30 percent (see Table 8-3 and Table 8-4). This
39 decreasing trend can be attributed to a 21 percent reduction in the amount of decomposable materials (i.e., paper and
40 paperboard, food scraps, and yard trimmings) discarded in MSW landfills over the time series (EPA 2010) and an
41 increase in the amount of landfill gas collected and combusted (i.e., used for energy or flared), resulting in lower net
42 CH₄ emissions from MSW landfills.²⁴⁹ For instance, in 1990, approximately 954 Gg of CH₄ were recovered and
43 combusted from landfills, while in 2011, approximately 8,177 Gg of CH₄ were combusted, representing an average
44 annual increase in the quantity of CH₄ recovered and combusted from 1990 to 2011 of 11 percent (see Annex 3.13).
45 In 2011, an estimated 71 new landfill gas-to-energy (LFGTE) projects and 29 new flares began operation (EPA
46 2012). While the amount of landfill gas collected and combusted continues to increase every year, the rate of
47 increase in collection and combustion no longer exceeds the rate of additional CH₄ generation from the amount of
48 organic MSW landfilled as the U.S. population grows.

49 The total amount of MSW generated is expected to increase as the U.S. population continues to grow. The
50 percentage of waste landfilled, however, may decline due to increased recycling and composting practices.

²⁴⁹ Due to a lack of data specific to industrial waste landfills, landfill gas recovery is only estimated for MSW landfills.

1 Additionally, the quantity of recovered CH₄ that is either flared or used for energy purposes is expected to
 2 continually increase as a result of 1996 federal regulations that require large MSW landfills to collect and combust
 3 landfill gas (see 40 CFR Part 60, Subpart Cc 2005 and 40 CFR Part 60, Subpart WWW 2005), as well as voluntary
 4 programs that encourage CH₄ recovery and use such as EPA's Landfill Methane Outreach Program (LMOP), and
 5 federal and state incentives that promote renewable energy (e.g., tax credits, low interest loans, and Renewable
 6 Portfolio Standards).

7 Table 8-3: CH₄ Emissions from Landfills (Tg CO₂ Eq.)

Activity	1990	2005	2007	2008	2009	2010	2011
MSW Landfills	172.6	241.2	254.2	259.2	262.9	266.6	270.2
Industrial Landfills	11.6	15.4	15.5	15.7	15.8	15.9	16.0
Recovered							
Gas-to-Energy	(13.3)	(55.9)	(62.6)	(67.2)	(74.2)	(82.5)	(88.0)
Flared	(6.7)	(75.7)	(83.2)	(81.5)	(78.6)	(81.4)	(83.7)
Oxidized ^a	(16.4)	(12.5)	(12.4)	(12.6)	(12.6)	(11.9)	(11.4)
Total	147.8	112.5	111.6	113.6	113.3	106.8	103.0

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

^a Includes oxidation at both municipal and industrial landfills. Oxidation at MSW landfills is accounted for after CH₄ recovery.

8 Table 8-4: CH₄ Emissions from Landfills (Gg)

Activity	1990	2005	2007	2008	2009	2010	2011
MSW Landfills	8,219	11,486	12,106	12,342	12,519	12,694	12,866
Industrial Landfills	554	733	740	746	752	758	761
Recovered							
Gas-to-Energy	(634)	(2,660)	(2,980)	(3,198)	(3,532)	(3,927)	(4,190)
Flared	(321)	(3,606)	(3,961)	(3,880)	(3,743)	(3,876)	(3,986)
Oxidized ^a	(782)	(595)	(590)	(601)	(600)	(565)	(545)
Total	7,037	5,357	5,314	5,409	5,397	5,084	4,906

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

^a Includes CH₄ oxidation at municipal and industrial landfills. Oxidation at MSW landfills is accounted for after CH₄ recovery.

9 Methodology

10 CH₄ emissions from landfills were estimated as the CH₄ produced from MSW landfills, plus the CH₄ produced by
 11 industrial waste landfills, minus the CH₄ recovered and combusted from MSW landfills, minus the CH₄ oxidized
 12 before being released into the atmosphere:

$$13 \quad \text{CH}_{4,\text{Solid Waste}} = [\text{CH}_{4,\text{MSW}} + \text{CH}_{4,\text{Ind}} - \text{R}] - \text{Ox}$$

14 where,

- 15 CH_{4,Solid Waste} = CH₄ emissions from solid waste
- 16 CH_{4,MSW} = CH₄ generation from MSW landfills,
- 17 CH_{4,Ind} = CH₄ generation from industrial landfills,
- 18 R = CH₄ recovered and combusted (only for MSW landfills), and
- 19 Ox = CH₄ oxidized from MSW and industrial waste landfills before release to the atmosphere.

20 The methodology for estimating CH₄ emissions from landfills is based on the first order decay model described by
 21 the IPCC (IPCC 2006). Methane generation is based on nationwide waste disposal data; it is not landfill-specific.
 22 The amount of CH₄ recovered, however, is landfill-specific, but only for MSW landfills due to a lack of data
 23 specific to industrial waste landfills. Values for the CH₄ generation potential (L₀) and decay rate constant (k) used in
 24 the first order decay model were obtained from an analysis of CH₄ recovery rates for a database of 52 landfills and
 25 from published studies of other landfills (RTI 2004; EPA 1998; SWANA 1998; Peer, Thorneloe, and Epperson
 26 1993). The decay rate constant was found to increase with average annual rainfall; consequently, values of k were
 27 developed for 3 ranges of rainfall, or climate types (wet, arid, and temperate). The annual quantity of waste placed
 28 in landfills was apportioned to the 3 ranges of rainfall based on the percent of the U.S. population in each of the 3

1 ranges. Historical census data were used to account for the shift in population to more arid areas over time. An
2 overview of the data sources and methodology used to calculate CH₄ generation and recovery is provided below,
3 while a more detailed description of the methodology used to estimate CH₄ emissions from landfills can be found in
4 Annex 3.13.

5 National MSW landfill waste generation and disposal data are obtained from the *BioCycle* State of Garbage surveys,
6 published approximately every two years. The State of Garbage (SOG) survey is the only continually updated
7 nationwide survey of waste disposed in landfills in the United States. The SOG surveys use the principles of mass
8 balance where all MSW generated is equal to the amount of MSW landfilled, combusted in waste-to-energy plants,
9 composted, and/or recycled (*BioCycle* 2010). This approach assumes that all waste management methods are
10 tracked and reported to state agencies. Survey respondents are asked to provide a breakdown of MSW generated
11 and managed by landfilling, recycling, composting, and combustion (in waste-to-energy facilities) in actual
12 tonnages. The survey reported data are adjusted to exclude non-MSW materials (e.g., industrial and agricultural
13 wastes, construction and demolition debris, automobile scrap, and sludge from wastewater treatment plants) that
14 may be included in survey responses. All state disposal data are adjusted for import/export; imported waste is
15 included in a particular state and exported waste is not. Where no waste generation data are provided by a state in
16 the SOG survey, the amount generated is estimated using the average nationwide waste per capita rate multiplied by
17 that particular state's population.

18 National landfill waste generation data for 1989 through 2008 were obtained from the SOG survey for every two
19 years (*BioCycle* 2006, 2008, and 2010). National landfill waste generation data for the years in-between the
20 *BioCycle* State of Garbage surveys (e.g., 2001, 2003, 2005, 2007, 2009, 2010, and 2011) were extrapolated based on
21 *BioCycle* data and the U.S. Census population. The most recent SOG survey was published in 2010 for the 2008
22 year. Waste generation data will be updated as new reports are published. Because the SOG survey does not
23 account for waste generated in U.S. territories, waste generation for the territories was estimated using population
24 data obtained from the U.S. Census Bureau (2012) and national per capita solid waste generation from the survey
25 (2010).

26 Estimates of the quantity of waste landfilled from 1989 to the current inventory year are determined by applying a
27 waste disposal factor to the total amount of waste generated (i.e., the SOG data). A waste disposal factor is
28 determined for each year an SOG survey is published and equals the ratio of the total amount of waste landfilled to
29 the total amount of waste generated. The waste disposal factor is interpolated for the years in-between the *BioCycle*
30 surveys, as is done for the amount of waste generated for a given survey year.

31 Estimates of the annual quantity of waste landfilled for 1960 through 1988 were obtained from EPA's
32 *Anthropogenic Methane Emissions in the United States, Estimates for 1990: Report to Congress* (EPA 1993) and an
33 extensive landfill survey by the EPA's Office of Solid Waste in 1986 (EPA 1988). Although waste placed in
34 landfills in the 1940s and 1950s contributes very little to current CH₄ generation, estimates for those years were
35 included in the first order decay model for completeness in accounting for CH₄ generation rates and are based on the
36 population in those years and the per capita rate for land disposal for the 1960s. For calculations in this inventory,
37 wastes landfilled prior to 1980 were broken into two groups: wastes disposed in landfills (Methane Conversion
38 Factor, MCF, of 1) and those disposed in dumps (MCF of 0.6). All calculations after 1980 assume waste is disposed
39 in managed, modern landfills. Please see Annex 3.13 for more details.

40 Methane recovery is currently only accounted for at MSW landfills since no comprehensive data regarding gas
41 collection systems have been published for industrial waste landfills. The estimated landfill gas recovered per year at
42 MSW landfills was based on a combination of three databases: the flare vendor database (contains updated sales
43 data collected from vendors of flaring equipment), a database of landfill gas-to-energy (LFGTE) projects compiled
44 by LMOP (EPA 2012), and a database developed by the Energy Information Administration (EIA) for the voluntary
45 reporting of greenhouse gases (EIA 2007). Based on the information provided by the EIA and flare vendor
46 databases, the CH₄ combusted by flares in operation from 1990 to the current inventory year was estimated.
47 Information provided by the EIA and LMOP databases were used to estimate CH₄ combusted in LFGTE projects
48 over the time series. The three databases were carefully compared to identify landfills that were in two or all three
49 of the databases to avoid double or triple counting CH₄ reductions.

50 The flare vendor database estimates CH₄ combusted by flares using the midpoint of a flare's reported capacity while
51 the EIA database uses landfill-specific measured gas flow. As the EIA database only includes data through 2006;
52 2007 to 2011 recovery for projects included in the EIA database were assumed to be the same as in 2006. This
53 quantity likely underestimates flaring because these databases do not have information on all flares in operation.

1 The EIA database is no longer being updated and it is expected that data obtained from the EPA's Greenhouse Gas
2 Reporting Program (GHGRP) will serve as a supplemental data source for facility-reported recovery data.
3 Additionally, the EIA and LMOP databases provided data on landfill gas flow and energy generation for landfills
4 with LFGTE projects. If a landfill in the EIA database was also in the LMOP and/or the flare vendor database, the
5 emissions avoided were based on the EIA data because landfill owners or operators reported the amount recovered
6 based on measurements of gas flow and concentration, and the reporting accounted for changes over time. If both
7 flare data and LMOP recovery data were available for any of the remaining landfills (i.e., not in the EIA database),
8 then the emissions recovery was based on the LMOP data, which provides reported landfill-specific data on gas flow
9 for direct use projects and project capacity (i.e., megawatts) for electricity projects. The flare data, on the other
10 hand, only provide a range of landfill gas flow for a given flare size. Given that each LFGTE project is likely to also
11 have a flare, double counting reductions from flares and LFGTE projects in the LMOP database was avoided by
12 subtracting emission reductions associated with LFGTE projects for which a flare had not been identified from the
13 emission reductions associated with flares (referred to as the flare correction factor). A further explanation of the
14 methodology used to estimate the landfill gas recovered can be found in Annex 3.13.

15 A destruction efficiency of 99 percent was applied to CH₄ recovered to estimate CH₄ emissions avoided due to the
16 combusting of CH₄ in destruction devices, i.e., flares. The destruction efficiency value was selected based on the
17 range of efficiencies (86 to 99 percent) recommended for flares in EPA's AP-42 Compilation of Air Pollutant
18 Emission Factors, Chapter 2.4 (EPA 2008), efficiencies used to establish new source performance standards (NSPS)
19 for landfills, and in recommendations for shutdown flares used in LMOP.

20 Emissions from industrial waste landfills were estimated from industrial production data (ERG 2012), waste
21 disposal factors, and the first order decay model. As over 99 percent of the organic waste placed in industrial waste
22 landfills originated from the food processing (meat, vegetables, fruits) and pulp and paper industries, estimates of
23 industrial landfill emissions focused on these two sectors (EPA 1993). There are currently no data sources that track
24 and report the amount and type of waste disposed of in industrial waste landfills in the United States. Therefore, the
25 amount of waste landfilled is assumed to be a fraction of production that is held constant over the time series as
26 explained in Annex 3.13. The composition of waste disposed of in industrial waste landfills is expected to be more
27 consistent in terms of composition and quantity than that disposed of in MSW landfills.

28 The amount of CH₄ oxidized by the landfill cover at both municipal and industrial waste landfills was assumed to be
29 ten percent of the CH₄ generated that is not recovered (IPCC 2006, Mancinelli and McKay 1985, Czepiel et al.
30 1996). To calculate net CH₄ emissions, both CH₄ recovered and CH₄ oxidized were subtracted from CH₄ generated
31 at municipal and industrial waste landfills.

32 Uncertainty and Time-Series Consistency

33 Several types of uncertainty are associated with the estimates of CH₄ emissions from MSW and industrial waste
34 landfills. The primary uncertainty concerns the characterization of landfills. Information is not available on two
35 fundamental factors affecting CH₄ production: the amount and composition of waste placed in every MSW and
36 industrial waste landfill for each year of its operation. The SOG survey is the only nationwide data source that
37 compiles the amount of MSW disposed at the state-level. The surveys do not include information on waste
38 composition and there are no comprehensive data sets that compile quantities of waste disposed or waste
39 composition by landfill. Some MSW landfills have conducted detailed waste composition studies, but landfills in
40 the United States are not required to perform these types of studies. The approach used here assumes that the CH₄
41 generation potential and the rate of decay that produces CH₄, as determined from several studies of CH₄ recovery at
42 MSW landfills, are representative of conditions at U.S. landfills. When this top-down approach is applied at the
43 nationwide level, the uncertainties are assumed to be less than when applying this approach to individual landfills
44 and then aggregating the results to the national level. In other words, this approach may over- and under-estimate
45 CH₄ generation at some landfills if used at the facility-level, but the end result is expected to balance out because it
46 is being applied nationwide. There is also a high degree of uncertainty and variability associated with the first order
47 decay model, particularly when a homogeneous waste composition and hypothetical decomposition rates are applied
48 to heterogeneous landfills (IPCC 2006).

49 Additionally, there is a lack of landfill-specific information regarding the number and type of industrial waste
50 landfills in the United States. The approach used here assumes that the majority (99 percent) of industrial waste
51 disposed of in industrial waste landfills consists of waste from the pulp and paper and food and beverage industries.
52 However, because waste generation and disposal data are not available in an existing data source for all U.S.

1 industrial waste landfills, we apply a straight disposal factor over the entire time series to the amount of waste
 2 generated to determine the amounts disposed.

3 Aside from the uncertainty in estimating CH₄ generation potential, uncertainty exists in the estimates of the landfill
 4 gas oxidized. A constant oxidation factor of 10 percent as recommended by the Intergovernmental Panel on Climate
 5 Change (IPCC) for managed landfills is used for both MSW and industrial waste landfills regardless of climate, the
 6 type of cover material, and/or presence of a gas collection system. The number of field studies measuring the rate of
 7 oxidation has increased substantially since the IPCC 2006 Guidelines were published and, as discussed in the
 8 Potential Improvements section, efforts are being made to review the literature and revise this value based on recent,
 9 peer-reviewed studies.

10 Another significant source of uncertainty lies with the estimates of CH₄ that are recovered by flaring and gas-to-
 11 energy projects at MSW landfills. Three separate databases containing recovery information are used to determine
 12 the total amount of CH₄ recovered and there are uncertainties associated with each. The LMOP database and the
 13 flare vendor databases are updated annually, while the EIA database has not been updated since 2005 and will
 14 essentially be replaced by the GHGRP data for a portion of landfills (i.e., those meeting the GHGRP thresholds). To
 15 avoid double counting and to use the most relevant estimate of CH₄ recovery for a given landfill, a hierarchical
 16 approach is used among the three databases. The EIA data are given precedence because CH₄ recovery was directly
 17 reported by landfills, the LMOP data are given second priority because CH₄ recovery is estimated from facility-
 18 reported LFGTE system characteristics, and the flare data are given third priority because this database contains
 19 minimal information about the flare and no site-specific operating characteristics (Bronstein et al., 2012). The IPCC
 20 default value of 10 percent for uncertainty in recovery estimates was used in the uncertainty analysis when metering
 21 of landfill gas was in place (for about 64 percent of the CH₄ estimated to be recovered). This 10 percent uncertainty
 22 factor applies to 2 of the 3 databases (EIA and LMOP). For flaring without metered recovery data (approximately 34
 23 percent of the CH₄ estimated to be recovered), a much higher uncertainty of approximately 50 percent was used
 24 (e.g., when recovery was estimated as 50 percent of the flare’s design capacity). The compounding uncertainties
 25 associated with the 3 databases leads to the large upper and lower bounds for MSW landfills presented in Table 8-5.

26 The results of the IPCC Good Practice Guidance Tier 2 quantitative uncertainty analysis are summarized in Table
 27 8-5. In 2011, landfill CH₄ emissions were estimated to be between 46.4 and 149.6 Tg CO₂ Eq., which indicates a
 28 range of 55 percent below to 45 percent above the 2011 emission estimate of 103.0 Tg CO₂ Eq.

29 **Table 8-5: Tier 2 Quantitative Uncertainty Estimates for CH₄ Emissions from Landfills (Tg CO₂ Eq. and Percent)**

Source	Gas	2011 Emission	Uncertainty Range Relative to Emission Estimate ^a			
		Estimate	(%)			
		(Tg CO ₂ Eq.)	Lower Bound	Upper Bound	Lower Bound	Upper Bound
Landfills	CH₄	103.0	46.4	149.6	-55%	+45%
MSW	CH ₄	88.6	33.5	134.3	-62%	+51%
Industrial	CH ₄	14.4	10.4	17.3	-28%	+20%

^a Range of emission estimates predicted by Monte Carlo Stochastic Simulation for a 95 percent confidence interval.

30 QA/QC and Verification

31 A QA/QC analysis was performed for data gathering and input, documentation, and calculation. QA/QC checks are
 32 not performed on the published data used to populate the Inventory data set, including the SOG survey data and the
 33 published LMOP database. A primary focus of the QA/QC checks was to ensure that CH₄ recovery estimates were
 34 not double-counted and that all LFGTE projects and flares were included in the respective project databases. Both
 35 manual and electronic checks were made to ensure that emission avoidance from each landfill was calculated in only
 36 one of the three databases. The primary calculation spreadsheet is tailored from the IPCC waste model and has been
 37 verified previously using the original, peer-reviewed IPCC waste model. All model input values were verified by
 38 secondary QA/QC review.

39 Recalculations Discussion

40 When conducted, methodological recalculations are applied to the entire time-series to ensure time-series
 41 consistency from 1990 through the current inventory year. No methodological changes were made for this

1 Inventory, but the national landfill waste generation data for 2007, 2008, 2009, and 2010 were recalculated for states
2 that did not report an amount of waste generated in the SOG 2010 survey. This recalculation was warranted after
3 reviewing the waste generation and disposal trends over the time series, particularly for years after 2004 where a
4 noticeable decrease in the amount of waste generated was calculated. For states that did not report an amount of
5 waste generated in the 2010 survey (*BioCycle 2010*), the recalculations used the most recent SOG waste per capita
6 data in the 2010 survey and state-specific generation rates from the previous SOG survey (*BioCycle 2008*). These
7 recalculations resulted in a slight increase in the waste generated for 2007 through 2010.

8 Planned Improvements

9 Improvements to the Inventory being examined include incorporating data from the EPA's GHGRP and recent peer-
10 reviewed literature, modifying the default oxidation factor applied to MSW and industrial waste landfills, and either
11 modifying the bulk waste degradable organic carbon (DOC) value or estimating emissions using a waste-specific
12 approach in the first order decay model.

13 Beginning in 2011, all MSW landfills that accepted waste on or after January 1, 1980 and generate CH₄ in amounts
14 equivalent to 25,000 metric tons or more of carbon dioxide equivalent (CO₂ Eq.) were required to calculate and
15 report their greenhouse gas emissions to EPA through its GHGRP. The MSW landfill source category of the
16 GHGRP consists of the landfill, landfill gas collection systems, and landfill gas destruction devices, including flares.
17 Potential improvements to the inventory methodology may be made using the GHGRP data, specifically for inputs
18 to the first order decay equation. The approach used by the inventory to estimate CH₄ generation assumes a bulk
19 waste-specific DOC value that may not accurately capture the changing waste composition over the time series (e.g.,
20 the reduction of organics entering the landfill environment due to increased composting, see Box 8-4). Using data
21 obtained from the GHGRP and any publicly available landfill-specific waste characterization studies in the United
22 States, the methodology may be modified to incorporate a waste composition approach or revisions may be made to
23 the bulk waste DOC value currently used. Additionally, GHGRP data could be analyzed and a weighted average for
24 the methane correction factor (MCF), fraction of CH₄ (F) in the landfill gas, the destruction efficiency of flares, and
25 the decay rate constant (k) could replace the values currently used in the inventory.

26 The most significant contribution of the GHGRP data to the Inventory is expected to be the amount of recovered
27 landfill gas and other information related to the gas collection system (Bronstein et al., 2012). Information for
28 landfills with gas collection systems reporting under the GHGRP will be incorporated into the inventory data set and
29 the measured CH₄ recovery data will be used for the reporting landfills in lieu of the EIA, LMOP, and flare vendor
30 data. The GHGRP data undergo an extensive series of verification steps, are more reliable and accurate than the
31 data currently used, and will reduce uncertainties surrounding CH₄ recovery when applied to the landfills in the
32 inventory data set (Bronstein et al., 2012).

33 In addition to MSW landfills, industrial waste landfills at facilities generating CH₄ in amounts equivalent to 25,000
34 metric tons or more of CO₂ Eq. were required to report their GHG emissions beginning in September 2012 through
35 EPA's GHGRP. Similar data for industrial waste landfills as is required for the MSW landfills will be reported. Any
36 additions or improvements to the inventory using reported GHGRP data will be made for the industrial waste
37 landfill portion of the inventory. One possible improvement is the addition of industrial sectors other than pulp and
38 paper, and food and beverage (e.g., metal foundries, petroleum refineries, and chemical manufacturing facilities).
39 Of particular interest in the GHGRP data set for industrial waste landfills will be the presence of gas collection
40 systems since recovery is not currently associated with industrial waste landfills in the inventory methodology. It is
41 unlikely that data reported through the GHGRP for industrial waste landfills will yield improved estimates for k and
42 L_o for the industrial sectors. However, EPA is considering an update to the L_o and k values for the pulp and paper
43 sector and are currently gathering feedback from stakeholders.

44 The addition of this higher tier data will improve the emission calculations to provide a more accurate representation
45 of greenhouse gas emissions from MSW and industrial waste landfills, but potential improvements to the Inventory
46 will not occur until after the deferral of GHGRP equation inputs expires in March 2013 for both MSW and industrial
47 waste landfills, or as early as the 1990 to 2013 Inventory report. Facility-level reporting data from the GHGRP are
48 not available for all Inventory years as reported in this Inventory; therefore, particular attention will be made to
49 ensure time series consistency while incorporating data from EPA's GHGRP that would be useful to improve the

1 emissions estimates for MSW landfills. In implementing improvements and integration of data from the GHGRP,
2 the latest guidance from the IPCC on the use of facility-level data in national inventories will be relied upon.²⁵⁰

3 As a first step toward revising the oxidation factor used in the Inventory, a literature review was conducted in 2011
4 (RTI 2011). A standard CH₄ oxidation factor of 10 percent has been used for both industrial and MSW landfills for
5 all Inventory reports and is currently recommended as the default for well-managed landfills in the latest IPCC
6 guidelines (2006). Recent comments on the Inventory methodology indicated that a default oxidation factor of 10
7 percent may be less than oxidation rates achieved at well-managed landfills with gas collection and control. The
8 impact of different landfill cover types on the rate of oxidation warrants further investigation as well.

9 Currently, one oxidation factor (10 percent) is applied to the total amount of waste generated nationwide. Changing
10 the oxidation factor and calculating the amount of CH₄ oxidized from landfills with gas collection and control
11 requires the estimation of waste disposed of in these types of landfills. The Inventory methodology uses waste
12 generation data from the SOG surveys, which report the total amount of waste generated and disposed nationwide
13 by state. In 2010, the State of Garbage survey requested data on the presence of landfill gas collection systems for
14 the first time. Twenty-eight states reported that 260 out of 1,414 (18 percent) operational landfills recovered landfill
15 gas (*BioCycle* 2010). However, the survey did not include closed landfills with gas collection and control systems.
16 In the future, the amount of states collecting and reporting this information is expected to increase. The EPA's
17 GHGRP data set for MSW landfills could be used to fill in the gaps related to the amount of waste disposed in
18 landfills with gas collection systems. Although the EPA's GHGRP does not capture every landfill in the United
19 States, larger landfills are expected to meet the reporting thresholds and will be reporting waste disposal information
20 by year beginning in March 2013. After incorporating the EPA's GHGRP data, it may be possible to calculate the
21 amount of waste disposed of at landfills with and without gas collection systems in the United States, which will
22 allow the Inventory waste model to apply different oxidation factors depending on the presence of a gas collection
23 system.

24 While research findings indicate some evidence that landfills with gas collection and control achieve a 20 percent or
25 higher oxidation rate, there is not sufficient certainty to adopt a higher oxidation rate at this time. It is expected that
26 with increased reporting by states in the State of Garbage survey, as well as the data collected through EPA's
27 GHGRP, the oxidation rate for at least a subset of landfills may be increased in a future Inventory. A continued
28 effort will be made to review peer-reviewed field studies that focus on oxidation specifically to determine how
29 oxidation is affected by the presence of a gas collection system and landfill cover type and whether increasing the
30 oxidation factor is warranted for all or only a portion of landfills (e.g., open versus closed, or only those with gas
31 collection systems).

32
33 [Begin Text Box]

34 Box 8-3: Nationwide Municipal Solid Waste Data Sources

35 Municipal solid waste generated in the United States can be managed through landfilling, recycling, composting,
36 and combustion with energy recovery. There are two main sources for nationwide solid waste management data in
37 the United States,

- 38 • The BioCycle and Earth Engineering Center of Columbia University's State of Garbage (SOG) in America
39 surveys and
- 40 • The EPA's Municipal Solid Waste in The United States: Facts and Figures reports.

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

²⁵⁰ See: http://www.ipcc-nggip.iges.or.jp/meeting/pdfiles/1008_Model_and_Facility_Level_Data_Report.pdf

1 The EPA Facts and Figures reports use a materials flow methodology, which relies heavily on a mass balance
2 approach. Data are gathered from industry associations, key businesses, similar industry sources, and government
3 agencies (e.g., the Department of Commerce and the U.S. Census Bureau) and are used to estimate tons of materials
4 and products generated, recycled, or discarded nationwide. The amount of MSW generated is estimated by
5 adjusting the imports and exports of produced materials. MSW that is not recycled, composted, or combusted is
6 assumed to be landfilled. The data presented in the report are nationwide totals.

7 The State of Garbage surveys are the preferred data source for estimating waste generation and disposal amounts in
8 the inventory because they are considered a more objective, numbers-based analysis of solid waste management in
9 the United States. However, the EPA Facts and Figures reports are useful when investigating waste management
10 trends at the nationwide level and for typical waste composition data, which the State of Garbage surveys do not ask
11 for.

12 In this Inventory, emissions from solid waste management are presented separately by waste management option,
13 except for recycling of waste materials. Emissions from recycling are attributed to the stationary combustion of
14 fossil fuels, and are presented in the stationary combustion chapter in the Energy sector, although the emissions
15 estimates are not called out separately. Emissions from solid waste disposal in landfills and the composting of solid
16 waste materials are presented in the Landfills and Composting chapters in the Waste sector of this report. In the
17 United States, almost all incineration of MSW occurs at waste-to-energy facilities or industrial facilities where
18 useful energy is recovered, and thus emissions from waste incineration are accounted for in the Incineration chapter
19 of the Energy sector of this report.

20

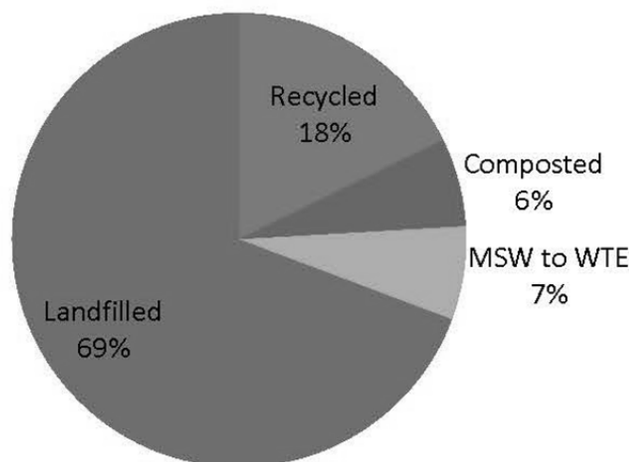
21 Box 8-4: Overview of the Waste Sector

22 As shown in Figure 8-2 and Figure 8-3, landfilling of MSW is currently and has been the most common waste
23 management practice. A large portion of materials in the waste stream are recovered for recycling and composting,
24 which is becoming an increasingly prevalent trend throughout the country. Materials that are composted would have
25 normally been disposed of in a landfill.

26

27 Figure 8-2: Management of Municipal Solid Waste in the United States, 2010 (BioCycle 2010)

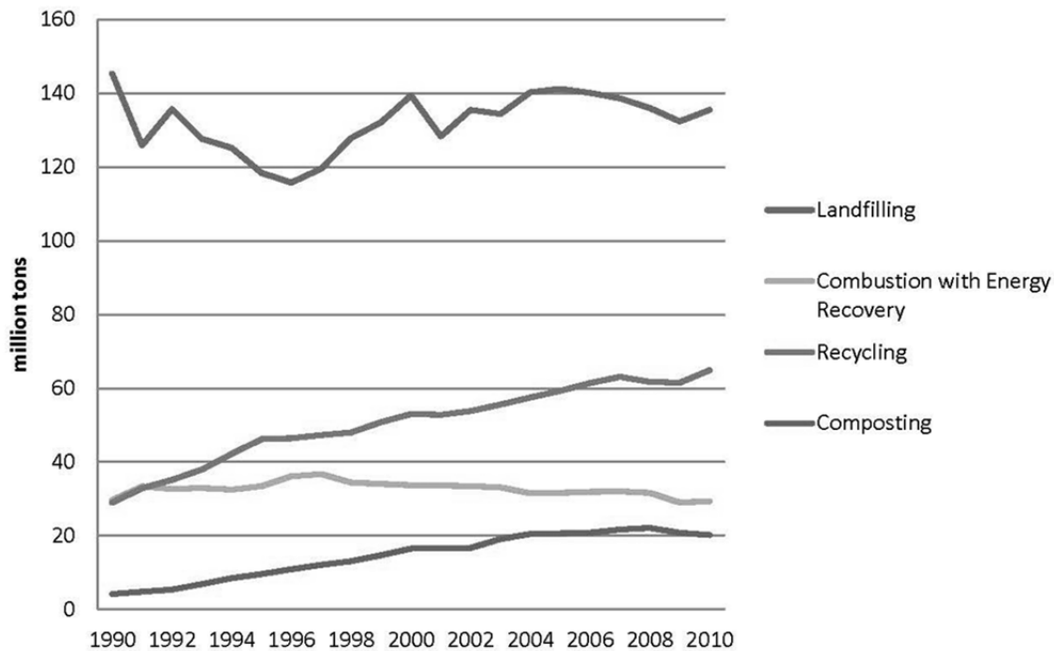
Management of MSW in the United States (BioCycle 2010)



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1 Figure 8-3: MSW Management Trends from 1990 to 2010 (EPA 2011)



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Table 8-6 presents a typical composition of waste disposed of at a typical MSW landfill in the United States over time. It is important to note that the actual composition of waste entering each landfill will vary from that presented in Table 8-6. Understanding how the waste composition changes over time, specifically for the degradable waste types, is important for estimating greenhouse gas emissions. For certain degradable waste types (i.e., paper and paperboard), the amounts discarded have decreased over time due to an increase in recovery (see Table 8-6 and Figure 8-4). Landfill ban legislation affecting yard trimmings resulted in an increase of composting from 1990 to 2008. Table 8-6 and Figure 8-4 do not reflect the impact of backyard composting on yard trimming generation and recovery estimates. The recovery of food trimmings has been consistently low. Increased recovery of degradable materials reduces the CH₄ generation potential and CH₄ emissions from landfills,

1 Table 8-6: Materials Discarded in the Municipal Waste Stream by Waste Type, percent (EPA 2011)

Waste Type	1990	2005	2007	2008	2009	2010
Paper and Paperboard	24.5%	24.5%	21.7%	19.7%	14.8%	15.3%
Glass	5.7%	5.7%	5.5%	5.3%	5.0%	4.8%
Metals	7.7%	7.7%	7.9%	8.0%	8.0%	8.3%
Plastics	15.7%	15.7%	16.4%	16.0%	15.8%	16.3%
Rubber and Leather	3.5%	3.5%	3.6%	3.7%	3.7%	3.8%
Textiles	5.5%	5.5%	5.9%	6.2%	6.3%	6.4%
Wood	7.4%	7.4%	7.5%	7.6%	7.7%	7.8%
Other ^a	1.8%	1.8%	1.9%	1.9%	1.9%	1.9%
Food Scraps ^b	17.9%	17.9%	18.2%	18.6%	19.1%	19.3%
Yard Trimmings ^c	7.0%	7.0%	6.7%	6.6%	7.6%	8.1%
Miscellaneous Inorganic Wastes	2.1%	2.1%	2.1%	2.2%	2.2%	2.2%

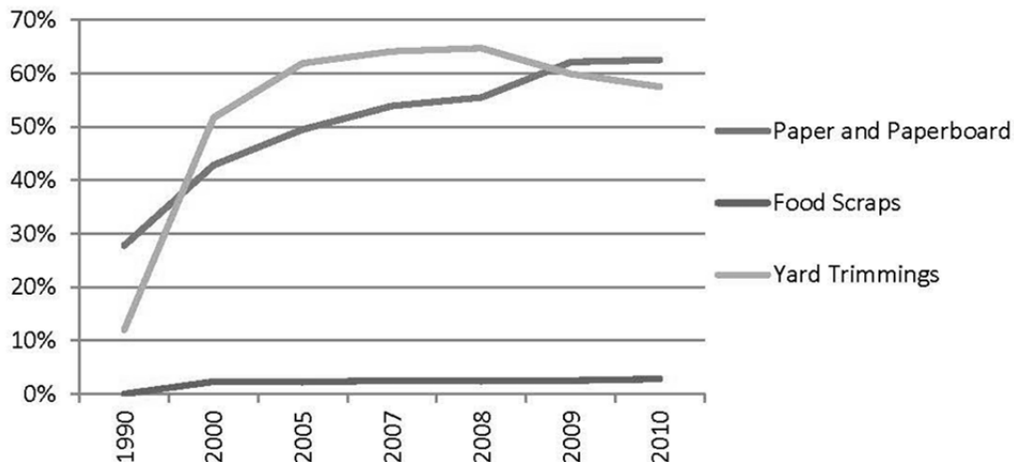
^a Includes electrolytes in batteries and fluff pulp, feces, and urine in disposable diapers. Details may not add to totals due to rounding. Source: EPA 2011.

^b Data for food scraps were estimated using sampling studies in various parts of the country in combination with demographic data on population, grocery store sales, restaurant sales, number of employees, and number of prisoners, students, and patients in institutions. Source: EPA 2010.

^c Data for yard trimmings were estimated using sampling studies, population data, and published sources documenting legislation affecting yard trimmings disposal in landfills. Source: EPA 2010.

2

3 Figure 8-4: Percent of Recovered Degradable Materials from 1990 to 2010, percent (EPA 2011)



4

5 [End Box]

6

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8 [Begin Text Box]

9 **Box 8-5: Description of a Modern, Managed Landfill**

10 Modern, managed landfills are well-engineered facilities that are located, designed, operated, and monitored to
 11 ensure compliance with federal, state, and tribal regulations. Municipal solid waste (MSW) landfills must be
 12 designed to protect the environment from contaminants which may be present in the solid waste stream.

13 Requirements for affected MSW landfills may include:

- 14 • Siting requirements to protect sensitive areas (e.g., airports, floodplains, wetlands, fault areas, seismic

1 impact zones, and unstable areas)

- 2 • Design requirements for new landfills to ensure that Maximum Contaminant Levels (MCLs) will not be
- 3 exceeded in the uppermost aquifer (e.g., composite liners and leachate collection systems)
- 4 • Leachate collection and removal systems
- 5 • Operating practices (e.g., daily and intermediate cover, receipt of regulated hazardous wastes, use of
- 6 landfill cover material, access options to prevent illegal dumping, use of a collection system to prevent
- 7 stormwater run-on/run-off, record-keeping)
- 8 • Air monitoring requirements (explosive gases)
- 9 • Groundwater monitoring requirements
- 10 • Closure and post-closure care requirements (e.g., final cover construction), and
- 11 • Corrective action provisions.

12 Specific federal regulations that affected MSW landfills must comply with include the 40 CFR Part 258 (Subtitle D
13 of RCRA), or equivalent state regulations and the New Source Performance Standards (NSPS) 40 CFR Part 60
14 Subpart WWW. Additionally, state and tribal requirements may exist. For more information regarding federal
15 MSW landfill regulations, see http://www.epa.gov/osw/nonhaz/municipal/landfill/msw_regs.htm.

16 [End Box]

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19 [Begin Text Box]

20 Box 8-6: Biogenic Wastes in Landfills

21 Regarding the depositing of wastes of biogenic origin in landfills (i.e., all degradable waste), empirical evidence
22 shows that some of these wastes degrade very slowly in landfills, and the C they contain is effectively sequestered in
23 landfills over a period of time (Barlaz 1998, 2006). Estimates of C removals from landfilling of forest products,
24 yard trimmings, and food scraps are further described in the Land Use, Land-Use Change, and Forestry chapter,
25 based on methods presented in IPCC (2003) and IPCC (2006).

26 [End Box]

28 **8.2. Wastewater Treatment (IPCC Source Category 6B)**

29 Wastewater treatment processes can produce anthropogenic CH₄ and N₂O emissions. Wastewater from domestic²⁵¹
30 and industrial sources is treated to remove soluble organic matter, suspended solids, pathogenic organisms, and
31 chemical contaminants. Treatment may either occur on site, most commonly through septic systems or package
32 plants, or off site at centralized treatment systems. Centralized wastewater treatment systems may include a variety
33 of processes, ranging from lagooning to advanced tertiary treatment technology for removing nutrients. In the
34 United States, approximately 20 percent of domestic wastewater is treated in septic systems or other on-site systems,
35 while the rest is collected and treated centrally (U.S. Census Bureau 2011).

36 Soluble organic matter is generally removed using biological processes in which microorganisms consume the
37 organic matter for maintenance and growth. The resulting biomass (sludge) is removed from the effluent prior to
38 discharge to the receiving stream. Microorganisms can biodegrade soluble organic material in wastewater under
39 aerobic or anaerobic conditions, where the latter condition produces CH₄. During collection and treatment,
40 wastewater may be accidentally or deliberately managed under anaerobic conditions. In addition, the sludge may be
41 further biodegraded under aerobic or anaerobic conditions. The generation of N₂O may also result from the
42 treatment of domestic wastewater during both nitrification and denitrification of the N present, usually in the form of
43 urea, ammonia, and proteins. These compounds are converted to nitrate (NO₃) through the aerobic process of

²⁵¹ Throughout the Inventory, emissions from domestic wastewater also include any commercial and industrial wastewater collected and co-treated with domestic wastewater.

1 nitrification. Denitrification occurs under anoxic conditions (without free oxygen), and involves the biological
 2 conversion of nitrate into dinitrogen gas (N₂). N₂O can be an intermediate product of both processes, but has
 3 typically been associated with denitrification. Recent research suggests that higher emissions of N₂O may in fact
 4 originate from nitrification (Ahn et al. 2010).

5 The principal factor in determining the CH₄ generation potential of wastewater is the amount of degradable organic
 6 material in the wastewater. Common parameters used to measure the organic component of the wastewater are the
 7 Biochemical Oxygen Demand (BOD) and Chemical Oxygen Demand (COD). Under the same conditions,
 8 wastewater with higher COD (or BOD) concentrations will generally yield more CH₄ than wastewater with lower
 9 COD (or BOD) concentrations. BOD represents the amount of oxygen that would be required to completely
 10 consume the organic matter contained in the wastewater through aerobic decomposition processes, while COD
 11 measures the total material available for chemical oxidation (both biodegradable and non-biodegradable). Because
 12 BOD is an aerobic parameter, it is preferable to use COD to estimate CH₄ production. The principal factor in
 13 determining the N₂O generation potential of wastewater is the amount of N in the wastewater. The variability of N
 14 in the influent to the treatment system, as well as the operating conditions of the treatment system itself, also impact
 15 the N₂O generation potential.

16 In 2011, CH₄ emissions from domestic wastewater treatment were 7.6Tg CO₂ Eq. (360 Gg). Emissions remained
 17 fairly steady from 1990 through 1997, but have decreased since that time due to decreasing percentages of
 18 wastewater being treated in anaerobic systems, including reduced use of on-site septic systems and central anaerobic
 19 treatment systems. In 2011, CH₄ emissions from industrial wastewater treatment were estimated to be 8.6 Tg CO₂
 20 Eq. (409 Gg). Industrial emission sources have increased across the time series through 1999 and then fluctuated up
 21 and down with production changes associated with the treatment of wastewater from the pulp and paper
 22 manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol production, and
 23 petroleum refining industries. Table 8-7 and Table 8-8 provide CH₄ and N₂O emission estimates from domestic and
 24 industrial wastewater treatment.

25 With respect to N₂O, the United States identifies two distinct sources for N₂O emissions from domestic wastewater:
 26 emissions from centralized wastewater treatment processes, and emissions from effluent from centralized treatment
 27 systems that has been discharged into aquatic environments. The 2011 emissions of N₂O from centralized
 28 wastewater treatment processes and from effluent were estimated to be 0.3 Tg CO₂ Eq. (1 Gg) and 4.9 Tg CO₂ Eq.
 29 (15.7 Gg), respectively. Total N₂O emissions from domestic wastewater were estimated to be 5.2 Tg CO₂ Eq. (16.7
 30 Gg). N₂O emissions from wastewater treatment processes gradually increased across the time series as a result of
 31 increasing U.S. population and protein consumption.

32
 33 Table 8-7: CH₄ and N₂O Emissions from Domestic and Industrial Wastewater Treatment (Tg CO₂ Eq.)

Activity	1990	2005	2007	2008	2009	2010	2011
CH₄	15.9	16.5	16.6	16.6	16.5	16.4	16.2
Domestic	8.8	8.3	8.1	8.0	8.0	7.8	7.6
Industrial*	7.1	8.2	8.5	8.6	8.5	8.6	8.6
N₂O	3.5	4.7	4.8	4.9	5.0	5.1	5.2
Domestic	3.5	4.7	4.8	4.9	5.0	5.1	5.2
Total	19.4	21.2	21.4	21.5	21.5	21.5	21.4

* Industrial activity includes the pulp and paper manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol production, and petroleum refining industries.

Note: Totals may not sum due to independent rounding.

34 Table 8-8: CH₄ and N₂O Emissions from Domestic and Industrial Wastewater Treatment (Gg)

Activity	1990	2005	2007	2008	2009	2010	2011
CH₄	758	785	791	791	786	779	770
Domestic	421	396	385	383	380	370	360
Industrial*	338	389	405	409	406	409	409
N₂O	11	15	16	16	16	16	17

* Industrial activity includes the pulp and paper manufacturing, meat and poultry processing, fruit and vegetable processing, starch-based ethanol production, and petroleum refining industries.

Note: Totals may not sum due to independent rounding.

1 Methodology

2 Domestic Wastewater CH₄ Emission Estimates

3 Domestic wastewater CH₄ emissions originate from both septic systems and from centralized treatment systems,
 4 such as publicly owned treatment works (POTWs). Within these centralized systems, CH₄ emissions can arise from
 5 aerobic systems that are not well managed or that are designed to have periods of anaerobic activity (e.g.,
 6 constructed wetlands), anaerobic systems (anaerobic lagoons and facultative lagoons), and from anaerobic digesters
 7 when the captured biogas is not completely combusted. CH₄ emissions from septic systems were estimated by
 8 multiplying the United States population by the percent of wastewater treated in septic systems (20 percent), an
 9 emission factor (10.7 g CH₄/capita/day) and converting that to Gg/year. Methane emissions from POTWs were
 10 estimated by multiplying the total BOD₅ produced in the United States by the percent of wastewater treated centrally
 11 (80 percent), the relative percentage of wastewater treated by aerobic and anaerobic systems, the relative percentage
 12 of wastewater facilities with primary treatment, the percentage of BOD₅ treated after primary treatment (67.5
 13 percent), the maximum CH₄-producing capacity of domestic wastewater (0.6), and the relative MCFs for aerobic
 14 (zero or 0.3) and anaerobic (0.8) systems with all aerobic systems assumed to be well-managed. Methane emissions
 15 from anaerobic digesters were estimated by multiplying the amount of biogas generated by wastewater sludge
 16 treated in anaerobic digesters by the proportion of CH₄ in digester biogas (0.65), the density of CH₄ (662 g CH₄/m³
 17 CH₄), and the destruction efficiency associated with burning the biogas in an energy/thermal device (0.99). The
 18 methodological equations are:

$$19 \quad \text{Emissions from Septic Systems} = A$$

$$20 \quad = US_{POP} \times (\% \text{ onsite}) \times (EF_{SEPTIC}) \times 1/10^9 \times \text{Days}$$

$$21 \quad \text{Emissions from Centrally Treated Aerobic Systems} = B$$

$$22 \quad = [(\% \text{ collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ aerobic}) \times (\% \text{ aerobic w/out primary}) + (\% \text{ collected}) \times (\text{total BOD}_5$$

$$23 \quad \text{produced}) \times (\% \text{ aerobic}) \times (\% \text{ aerobic w/primary}) \times (1 - \% \text{ BOD removed in prim. treat.})] \times (\% \text{ operations not well}$$

$$24 \quad \text{managed}) \times (B_o) \times (\text{MCF-aerobic_not_well_man}) \times 1/10^6$$

$$25 \quad \text{Emissions from Centrally Treated Anaerobic Systems} = C$$

$$26 \quad = [(\% \text{ collected}) \times (\text{total BOD}_5 \text{ produced}) \times (\% \text{ anaerobic}) \times (\% \text{ anaerobic w/out primary}) + (\% \text{ collected}) \times (\text{total}$$

$$27 \quad \text{BOD}_5 \text{ produced}) \times (\% \text{ anaerobic}) \times (\% \text{ anaerobic w/primary}) \times (1 - \% \text{ BOD removed in prim. treat.})] \times (B_o) \times (\text{MCF-}$$

$$28 \quad \text{anaerobic}) \times 1/10^6$$

$$29 \quad \text{Emissions from Anaerobic Digesters} = D$$

$$30 \quad = [(\text{POTW_flow_AD}) \times (\text{digester gas}) / (\text{per capita flow})] \times \text{conversion to m}^3 \times (\text{FRAC_CH}_4) \times (365.25) \times (\text{density}$$

$$31 \quad \text{of CH}_4) \times (1 - \text{DE}) \times 1/10^9$$

$$32 \quad \text{Total CH}_4 \text{ Emissions (Gg)} = A + B + C + D$$

33 where,

34	US _{POP}	= U.S. population
35	% onsite	= Flow to septic systems / total flow
36	% collected	= Flow to POTWs / total flow
37	% aerobic	= Flow to aerobic systems / total flow to POTWs
38	% anaerobic	= Flow to anaerobic systems / total flow to POTWs
39	% aerobic w/out primary	= Percent of aerobic systems that do not employ primary treatment
40	% aerobic w/primary	= Percent of aerobic systems that employ primary treatment
41	% BOD removed in prim. treat.	= 32.5%
42	% operations not well managed	= Percent of aerobic systems that are not well managed and in which
43		some anaerobic degradation occurs
44	% anaerobic w/out primary	= Percent of anaerobic systems that do not employ primary treatment

1	% anaerobic w/primary	= Percent of anaerobic systems that employ primary treatment
2	EF _{SEPTIC}	= Methane emission factor (10.7 g CH ₄ /capita/day) – septic systems
3	Days	= days per year (365.25)
4	Total BOD ₅ produced	= kg BOD/capita/day × U.S. population × 365.25 days/yr
5	B _o	= Maximum CH ₄ -producing capacity for domestic wastewater (0.60 kg CH ₄ /kg BOD)
6		
7	1/10 ⁶	= Conversion factor, kg to Gg
8	MCF-aerobic_not_well_man.	= CH ₄ correction factor for aerobic systems that are not well managed (0.3)
9		
10	MCF-anaerobic	= CH ₄ correction factor for anaerobic systems (0.8)
11	DE	= CH ₄ destruction efficiency from flaring or burning in engine (0.99 for enclosed flares)
12		
13	POTW_flow_AD	= Wastewater influent flow to POTWs that have anaerobic digesters (gal)
14	digester gas	= Cubic feet of digester gas produced per person per day (1.0 ft ³ /person/day) (Metcalf and Eddy 2003)
15		
16	per capita flow	= Wastewater flow to POTW per person per day (100 gal/person/day)
17	conversion to m ³	= Conversion factor, ft ³ to m ³ (0.0283)
18	FRAC_CH ₄	= Proportion CH ₄ in biogas (0.65)
19	density of CH ₄	= 662 (g CH ₄ /m ³ CH ₄)
20	1/10 ⁹	= Conversion factor, g to Gg

21 U.S. population data were taken from the U.S. Census Bureau International Database (U.S. Census 2012) and
 22 include the populations of the United States, American Samoa, Guam, Northern Mariana Islands, Puerto Rico, and
 23 the Virgin Islands. Table 8-9 presents U.S. population and total BOD₅ produced for 1990 through 2011, while Table
 24 8-10 presents domestic wastewater CH₄ emissions for both septic and centralized systems in 2011. The proportions
 25 of domestic wastewater treated onsite versus at centralized treatment plants were based on data from the 1989, 1991,
 26 1993, 1995, 1997, 1999, 2001, 2003, 2005, 2007, 2009, 2011 American Housing Surveys conducted by the U.S.
 27 Census Bureau (U.S. Census 2011), with data for intervening years obtained by linear interpolation. The percent of
 28 wastewater flow to aerobic and anaerobic systems, the percent of aerobic and anaerobic systems that do and do not
 29 employ primary treatment, and the wastewater flow to POTWs that have anaerobic digesters were obtained from the
 30 1992, 1996, 2000, and 2004 Clean Watershed Needs Survey (EPA 1992, 1996, 2000, and 2004). Data for
 31 intervening years were obtained by linear interpolation and the years 2004 through 2011 were forecasted from the
 32 rest of the time series. The BOD₅ production rate (0.09 kg/capita/day) and the percent BOD₅ removed by primary
 33 treatment for domestic wastewater were obtained from Metcalf and Eddy (2003). The CH₄ emission factor (0.6 kg
 34 CH₄/kg BOD₅) and the MCF used for centralized treatment systems were taken from IPCC (2006), while the CH₄
 35 emission factor (10.7 g CH₄/capita/day) used for septic systems were taken from Leverenz et al. (2010). The CH₄
 36 destruction efficiency for methane recovered from sludge digestion operations, 99 percent, was selected based on the
 37 range of efficiencies (98 to 100 percent) recommended for flares in AP-42 Compilation of Air Pollutant Emission
 38 Factors, Chapter 2.4 (EPA 1998), efficiencies used to establish new source performance standards (NSPS) for
 39 landfills, and in recommendations for closed flares used by the Landfill Methane Outreach Program (LMOP). The
 40 cubic feet of digester gas produced per person per day (1.0 ft³/person/day) and the proportion of CH₄ in biogas
 41 (0.65) come from Metcalf and Eddy (2003). The wastewater flow to a POTW (100 gal/person/day) was taken from
 42 the Great Lakes-Upper Mississippi River Board of State and Provincial Public Health and Environmental Managers,
 43 "Recommended Standards for Wastewater Facilities (Ten-State Standards)" (2004).

44 Table 8-9: U.S. Population (Millions) and Domestic Wastewater BOD₅ Produced (Gg)

Year	Population	BOD ₅
1990	253	8,333
2005	300	9,853
2007	305	10,039
2008	308	10,132
2009	311	10,220
2010	313	10,303
2011	316	10,377

Source: U.S. Census Bureau (2012);
Metcalf & Eddy 2003.

1 Table 8-10: Domestic Wastewater CH₄ Emissions from Septic and Centralized Systems (2011)

	CH ₄ emissions (Tg CO ₂ Eq.)	% of Domestic Wastewater CH ₄
Septic Systems	5.0	66.4%
Centralized Systems	2.5	33.6%
Total	7.6	100%

Note: Totals may not sum due to independent rounding.

2 **Industrial Wastewater CH₄ Emission Estimates**

3 Methane emissions estimates from industrial wastewater were developed according to the methodology described in
4 IPCC (2006). Industry categories that are likely to produce significant CH₄ emissions from wastewater treatment
5 were identified. High volumes of wastewater generated and a high organic wastewater load were the main criteria.
6 The top five industries that meet these criteria are pulp and paper manufacturing; meat and poultry processing;
7 vegetables, fruits, and juices processing; starch-based ethanol production; and petroleum refining. Wastewater
8 treatment emissions for these sectors for 2011 are displayed in Table 8-11 below. Table 8-12 contains production
9 data for these industries.

10 Table 8-11: Industrial Wastewater CH₄ Emissions by Sector (2011)

	CH ₄ emissions (Tg CO ₂ Eq.)	% of Industrial Wastewater CH ₄
Pulp & Paper	4.1	48%
Meat & Poultry	3.7	43%
Petroleum Refineries	0.6	7%
Fruit & Vegetables	0.1	1%
Ethanol Refineries	0.1	1%
Total	8.6	100%

Note: Totals may not sum due to independent rounding.

11

12 Table 8-12: U.S. Pulp and Paper, Meat, Poultry, Vegetables, Fruits and Juices, Ethanol, and Petroleum Refining
13 Production (Tg)

Year	Pulp and Paper ^a	Meat (Live Weight Killed)	Poultry (Live Weight Killed)	Vegetables, Fruits and Juices	Ethanol	Petroleum Refining
1990	128.9	27.3	14.6	38.7	2.7	702.4
2005	131.4	31.4	25.1	42.9	11.7	818.6
2007	135.9	33.4	26.0	44.7	19.4	827.6
2008	134.5	34.4	26.6	45.1	26.9	836.8
2009	137.0	33.8	25.2	46.5	31.7	822.4
2010	137.0	33.7	25.9	43.2	39.5	848.6
2011	137.0	33.8	26.2	42.9	41.5	858.8

^aPulp and paper production is the sum of woodpulp production plus paper and paperboard production.

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 percentage of organic loading assumed to degrade
16 anaerobically, and the maximum CH₄ producing potential of industrial wastewater (B₀). Ratios of BOD:COD in
17 various industrial wastewaters were obtained from EPA (1997a) and used to estimate COD loadings. The B₀ value

1 used for all industries is the IPCC default value of 0.25 kg CH₄/kg COD (IPCC 2006).

2 For each industry, the percent of plants in the industry that treat wastewater on site, the percent of plants that have a
 3 primary treatment step prior to biological treatment, and the percent of plants that treat wastewater anaerobically
 4 were defined. The percent of wastewater treated anaerobically onsite (TA) was estimated for both primary treatment
 5 (%TA_p) and secondary treatment (%TA_s). For plants that have primary treatment in place, an estimate of COD that
 6 is removed prior to wastewater treatment in the anaerobic treatment units was incorporated.

7 The methodological equations are:

$$8 \quad \text{CH}_4 (\text{industrial wastewater}) = [P \times W \times \text{COD} \times \% \text{TA}_p \times B_o \times \text{MCF}] + [P \times W \times \text{COD} \times \% \text{TA}_s \times B_o \times \text{MCF}]$$

$$9 \quad \% \text{TA}_p = [\% \text{Plants}_o \times \% \text{WW}_{a,p} \times \% \text{COD}_p]$$

$$10 \quad \% \text{TA}_s = [\% \text{Plants}_a \times \% \text{WW}_{a,s} \times \% \text{COD}_s] + [\% \text{Plants}_t \times \% \text{WW}_{a,t} \times \% \text{COD}_s]$$

11 where,

12 CH₄ (industrial wastewater) = Total CH₄ emissions from industrial wastewater (kg/year)

13 P = Industry output (metric tons/year)

14 W = Wastewater generated (m³/metric ton of product)

15 COD = Organics loading in wastewater (kg/m³)

16 %TA_p = Percent of wastewater treated anaerobically on site in primary treatment

17 %TA_s = Percent of wastewater treated anaerobically on site in secondary treatment

18 %Plants_o = Percent of plants with onsite treatment

19 %WW_{a,p} = Percent of wastewater treated anaerobically in primary treatment

20 %COD_p = Percent of COD entering primary treatment

21 %Plants_a = Percent of plants with anaerobic secondary treatment

22 %Plants_t = Percent of plants with other secondary treatment

23 %WW_{a,s} = Percent of wastewater treated anaerobically in anaerobic secondary treatment

24 %WW_{a,t} = percent of wastewater treated anaerobically in other secondary treatment

25 %COD_s = percent of COD entering secondary treatment

26 B_o = Maximum CH₄ producing potential of industrial wastewater (default value of
 27 0.25 kg CH₄/kg COD)

28 MCF = CH₄ correction factor, indicating the extent to which the organic content
 29 (measured as COD) degrades anaerobically

30 As described below, the values presented in Table 8-13 were used in the emission calculations and are described in
 31 detail in Aguiar and Bartram (2008).

32 Table 8-13: Variables Used to Calculate Percent Wastewater Treated Anaerobically by Industry (%)

Variable	Industry						
	Pulp and Paper	Meat Processing	Poultry Processing	Fruit/Vegetable Processing	Ethanol Production – Wet Mill	Ethanol Production – Dry Mill	Petroleum Refining
%TA _p	0	0	0	0	0	0	0
%TA _s	10.5	33	25	4.2	33.3	75	100
%Plants _o	60	100	100	11	100	100	100
%Plants _a	25	33	25	5.5	33.3	75	100
%Plants _t	35	67	75	5.5	66.7	25	0
%WW _{a,p}	0	0	0	0	0	0	0
%WW _{a,s}	100	100	100	100	100	100	100
%WW _{a,t}	0	0	0	0	0	0	0
%COD _p	100	100	100	100	100	100	100
%COD _s	42	100	100	77	100	100	100

Source: Aguiar and Bartram (2008) Planned Revisions of the Industrial Wastewater Inventory Emission Estimates for the 1990-2007 Inventory. August 10, 2008.

33 *Pulp and Paper.* Wastewater treatment for the pulp and paper industry typically includes neutralization, screening,

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

12 Approximately 42 percent of the BOD passes on to secondary treatment, which consists of activated sludge, aerated
 13 stabilization basins, or non-aerated stabilization basins. No anaerobic activity is assumed to occur in activated
 14 sludge systems or aerated stabilization basins (note: although IPCC recognizes that some CH₄ can be emitted from
 15 anaerobic pockets, they recommend an MCF of zero). However, about 25 percent of the wastewater treatment
 16 systems used in the United States are non-aerated stabilization basins. These basins are typically 10 to 25 feet deep.
 17 These systems are classified as anaerobic deep lagoons (MCF = 0.8).

18 A time series of CH₄ emissions for 1990 through 2001 was developed based on production figures reported in the
 19 Lockwood-Post Directory (Lockwood-Post 2002). Published data from the American Forest and Paper Association,
 20 data published by Paper Loop, and other published statistics were used to estimate production for 2002 through 2011
 21 (Pulp and Paper 2005, 2006, and monthly reports from 2003 through 2008; Paper 360° 2007). The overall
 22 wastewater outflow was estimated to be 85 m³/metric ton, and the average BOD concentrations in raw wastewater
 23 was estimated to be 0.4 gram BOD/liter (EPA 1997b, EPA 1993, World Bank 1999). The COD:BOD ratio used to
 24 convert the organic loading to COD for pulp and paper facilities was 2 (EPA 1997a).

25 *Meat and Poultry Processing.* The meat and poultry processing industry makes extensive use of anaerobic lagoons
 26 in sequence with screening, fat traps and dissolved air flotation when treating wastewater on site. About 33 percent
 27 of meat processing operations (EPA 2002) and 25 percent of poultry processing operations (U.S. Poultry 2006)
 28 perform on-site treatment in anaerobic lagoons. The IPCC default B₀ of 0.25 kg CH₄/kg COD and default MCF of
 29 0.8 for anaerobic lagoons were used to estimate the CH₄ produced from these on-site treatment systems. Production
 30 data, in carcass weight and live weight killed for the meat and poultry industry, were obtained from the USDA
 31 Agricultural Statistics Database and the Agricultural Statistics Annual Reports (USDA 2012). Data collected by
 32 EPA's Office of Water provided estimates for wastewater flows into anaerobic lagoons: 5.3 and 12.5 m³/metric ton
 33 for meat and poultry production (live weight killed), respectively (EPA 2002). The loadings are 2.8 and 1.5 g
 34 BOD/liter for meat and poultry, respectively. The COD:BOD ratio used to convert the organic loading to COD for
 35 both meat and poultry facilities was 3 (EPA 1997a).

36 *Vegetables, Fruits, and Juices Processing.* Treatment of wastewater from fruits, vegetables, and juices processing
 37 includes screening, coagulation/settling, and biological treatment (lagooning). The flows are frequently seasonal,
 38 and robust treatment systems are preferred for on-site treatment. Effluent is suitable for discharge to the sewer.
 39 This industry is likely to use lagoons intended for aerobic operation, but the large seasonal loadings may develop
 40 limited anaerobic zones. In addition, some anaerobic lagoons may also be used (Nemerow and Dasgupta 1991).
 41 Consequently, 4.2 percent of these wastewater organics are assumed to degrade anaerobically. The IPCC default B₀
 42 of 0.25 kg CH₄/kg COD and default MCF of 0.8 for anaerobic treatment were used to estimate the CH₄ produced
 43 from these on-site treatment systems. The USDA National Agricultural Statistics Service (USDA 2012) provided
 44 production data for potatoes, other vegetables, citrus fruit, non-citrus fruit, and grapes processed for wine. Outflow
 45 and BOD data, presented in Table 8-14, were obtained from EPA (1974) for potato, citrus fruit, and apple
 46 processing, and from EPA (1975) for all other sectors. The COD:BOD ratio used to convert the organic loading to
 47 COD for all fruit, vegetable, and juice facilities was 1.5 (EPA 1997a).

48 Table 8-14: Wastewater Flow (m³/ton) and BOD Production (g/L) for U.S. Vegetables, Fruits, and Juices Production

Commodity	Wastewater Outflow (m ³ /ton)	BOD (g/L)
Vegetables		
Potatoes	10.27	1.765
Other Vegetables	8.69	0.794

Fruit

Apples	3.66	1.371
Citrus	10.11	0.317
Non-citrus	12.42	1.204
Grapes (for wine)	2.78	1.831

1 *Ethanol Production.* Ethanol, or ethyl alcohol, is produced primarily for use as a fuel component, but is also used in
 2 industrial applications and in the manufacture of beverage alcohol. Ethanol can be produced from the fermentation
 3 of sugar-based feedstocks (e.g., molasses and beets), starch- or grain-based feedstocks (e.g., corn, sorghum, and
 4 beverage waste), and cellulosic biomass feedstocks (e.g., agricultural wastes, wood, and bagasse). Ethanol can also
 5 be produced synthetically from ethylene or hydrogen and carbon monoxide. However, synthetic ethanol comprises
 6 only about 2 percent of ethanol production, and although the Department of Energy predicts cellulosic ethanol to
 7 greatly increase in the coming years, currently it is only in an experimental stage in the United States. According to
 8 the Renewable Fuels Association, 82 percent of ethanol production facilities use corn as the sole feedstock and 7
 9 percent of facilities use a combination of corn and another starch-based feedstock. The fermentation of corn is the
 10 principal ethanol production process in the United States and is expected to increase through 2012, and potentially
 11 more; therefore, emissions associated with wastewater treatment at starch-based ethanol production facilities were
 12 estimated (ERG 2006).

13 Ethanol is produced from corn (or other starch-based feedstocks) primarily by two methods: wet milling and dry
 14 milling. Historically, the majority of ethanol was produced by the wet milling process, but now the majority is
 15 produced by the dry milling process. The wastewater generated at ethanol production facilities is handled in a
 16 variety of ways. Dry milling facilities often combine the resulting evaporator condensate with other process
 17 wastewaters, such as equipment wash water, scrubber water, and boiler blowdown and anaerobically treat this
 18 wastewater using various types of digesters. Wet milling facilities often treat their steepwater condensate in
 19 anaerobic systems followed by aerobic polishing systems. Wet milling facilities may treat the stillage (or processed
 20 stillage) from the ethanol fermentation/distillation process separately or together with steepwater and/or wash water.
 21 CH₄ generated in anaerobic digesters is commonly collected and either flared or used as fuel in the ethanol
 22 production process (ERG 2006).

23 Available information was compiled from the industry on wastewater generation rates, which ranged from 1.25
 24 gallons per gallon ethanol produced (for dry milling) to 10 gallons per gallon ethanol produced (for wet milling)
 25 (Ruocco 2006a,b; Merrick 1998; Donovan 1996; and NRBP 2001). COD concentrations were also found to be
 26 about 3 g/L (Ruocco 2006a; Merrick 1998; White and Johnson 2003). The amount of wastewater treated
 27 anaerobically was estimated, along with how much of the CH₄ is recovered through the use of biomethanators (ERG
 28 2006). Methane emissions were then estimated as follows:

29

$$30 \text{ Methane} = [\text{Production} \times \text{Flow} \times \text{COD} \times 3.785 \times (\% \text{Plants}_o \times \% \text{WW}_{a,p} \times \% \text{COD}_p) + [\% \text{Plants}_a \times \% \text{WW}_{a,s} \times \% \text{COD}_s] +$$

$$31 [\% \text{Plants}_t \times \% \text{WW}_{a,t} \times \% \text{COD}_s]) \times B_o \times \text{MCF} \times \% \text{Not Recovered}] + [\text{Production} \times \text{Flow} \times 3.785 \times \text{COD} \times (\% \text{Plants}_o \times$$

$$32 \% \text{WW}_{a,p} \times \% \text{COD}_p) + [\% \text{Plants}_a \times \% \text{WW}_{a,s} \times \% \text{COD}_s] + [\% \text{Plants}_t \times \% \text{WW}_{a,t} \times \% \text{COD}_s]) \times B_o \times \text{MCF} \times (\% \text{Recovered}) \times$$

$$33 (1-DE)] \times 1/10^9$$

34 where,

- 35 Production = gallons ethanol produced (wet milling or dry milling)
- 36 Flow = gallons wastewater generated per gallon ethanol produced (1.25 dry milling, 10 wet milling)
- 37 COD = COD concentration in influent (3 g/l)
- 38 3.785 = conversion, gallons to liters
- 39 %Plants_o = percent of plants with onsite treatment (100%)
- 40 %WW_{a,p} = percent of wastewater treated anaerobically in primary treatment (0%)
- 41 %COD_p = percent of COD entering primary treatment (100%)
- 42 %Plants_a = percent of plants with anaerobic secondary treatment (33.3% wet, 75% dry)
- 43 %Plants_t = percent of plants with other secondary treatment (66.7% wet, 25% dry)
- 44 %WW_{a,s} = percent of wastewater treated anaerobically in anaerobic secondary treatment (100%)
- 45 %WW_{a,t} = percent of wastewater treated anaerobically in other secondary treatment (0%)
- 46 %COD_s = percent of COD entering secondary treatment (100%)
- 47 B_o = maximum methane producing capacity (0.25 g CH₄/g COD)

1	MCF	= methane conversion factor (0.8 for anaerobic systems)
2	% Recovered	= percent of wastewater treated in system with emission recovery
3	% Not Recovered	= 1 - percent of wastewater treated in system with emission recovery
4	DE	= destruction efficiency of recovery system (99%)
5	1/10 ⁹	= conversion factor, g to Gg

6 A time series of CH₄ emissions for 1990 through 2011 was developed based on production data from the Renewable
7 Fuels Association (RFA 2012).

8 *Petroleum Refining.* Petroleum refining wastewater treatment operations produce CH₄ emissions from anaerobic
9 wastewater treatment. The wastewater inventory section includes CH₄ emissions from petroleum refining
10 wastewater treated on site under intended or unintended anaerobic conditions. Most facilities use aerated biological
11 systems, such as trickling filters or rotating biological contactors; these systems can also exhibit anaerobic
12 conditions that can result in the production of CH₄. Oil/water separators are used as a primary treatment method;
13 however, it is unlikely that any COD is removed in this step.

14 Available information from the industry was compiled. The wastewater generation rate, from CARB (2007) and
15 Timm (1985), was determined to be 35 gallons per barrel of finished product. An average COD value in the
16 wastewater was estimated at 0.45 kg/m³ (Benyahia et al. 2006).

17 The equation used to calculate CH₄ generation at petroleum refining wastewater treatment systems is presented
18 below:

$$19 \text{ Methane} = \text{Flow} \times \text{COD} \times B_o \times \text{MCF}$$

20 where,

21	Flow	= Annual flow treated through anaerobic treatment system (m ³ /year)
22	COD	= COD loading in wastewater entering anaerobic treatment system (kg/m ³)
23	B _o	= maximum methane producing potential of industrial wastewater (default value of 0.25 24 kg CH ₄ /kg COD)
25	MCF	= methane conversion factor (0.3)

26
27 A time series of CH₄ emissions for 1990 through 2011 was developed based on production data from the Energy
28 Information Association (EIA 2012).

29 Domestic Wastewater N₂O Emission Estimates

30 N₂O emissions from domestic wastewater (wastewater treatment) were estimated using the IPCC (2006)
31 methodology, including calculations that take into account N removal with sewage sludge, non-consumption and
32 industrial/commercial wastewater N, and emissions from advanced centralized wastewater treatment plants:

- 33 • In the United States, a certain amount of N is removed with sewage sludge, which is applied to land, incinerated,
34 or landfilled (N_{SLUDGE}). The N disposal into aquatic environments is reduced to account for the sewage sludge
35 application.
- 36 • The IPCC methodology uses annual, per capita protein consumption (kg protein/[person-year]). For this
37 inventory, the amount of protein available to be consumed is estimated based on per capita annual food
38 availability data and its protein content, and then adjusts that data using a factor to account for the fraction of
39 protein actually consumed.
- 40 • Small amounts of gaseous nitrogen oxides are formed as byproducts in the conversion of nitrate to N gas in
41 anoxic biological treatment systems. Approximately 7 g N₂O is generated per capita per year if wastewater
42 treatment includes intentional nitrification and denitrification (Scheehle and Doorn 2001). Analysis of the 2004
43 CWNS shows that plants with denitrification as one of their unit operations serve a population of 2.4 million
44 people. Based on an emission factor of 7 g per capita per year, approximately 21.2 metric tons of additional N₂O
45 may have been emitted via denitrification in 2004. Similar analyses were completed for each year in the
46 Inventory using data from CWNS on the amount of wastewater in centralized systems treated in denitrification
47 units. Plants without intentional nitrification/denitrification are assumed to generate 3.2 g N₂O per capita per
48 year.

1 N₂O emissions from domestic wastewater were estimated using the following methodology:

$$\begin{aligned}
 2 \quad & N_2O_{TOTAL} = N_2O_{PLANT} + N_2O_{EFFLUENT} \\
 3 \quad & N_2O_{PLANT} = N_2O_{NIT/DENIT} + N_2O_{WOUT\ NIT/DENIT} \\
 4 \quad & N_2O_{NIT/DENIT} = [(US_{POPND}) \times EF_2 \times F_{IND-COM}] \times 1/10^9 \\
 5 \quad & N_2O_{WOUT\ NIT/DENIT} = \{[(US_{POP} \times WWTP) - US_{POPND}] \times F_{IND-COM} \times EF_1\} \times 1/10^9 \\
 6 \quad & N_2O_{EFFLUENT} = \{[\{((US_{POP} \times WWTP) - (0.9 \times US_{POPND})) \times Protein \times F_{NPR} \times F_{NON-CON} \times F_{IND-COM}) - N_{SLUDGE}\} \times EF_3 \times \\
 7 \quad & \quad \quad \quad 44/28\} \times 1/10^6
 \end{aligned}$$

8 where,

- 9 N₂O_{TOTAL} = Annual emissions of N₂O (Gg)
- 10 N₂O_{PLANT} = N₂O emissions from centralized wastewater treatment plants (Gg)
- 11 N₂O_{NIT/DENIT} = N₂O emissions from centralized wastewater treatment plants with
- 12 nitrification/denitrification (Gg)
- 13 N₂O_{WOUT NIT/DENIT} = N₂O emissions from centralized wastewater treatment plants without
- 14 nitrification/denitrification (Gg)
- 15 N₂O_{EFFLUENT} = N₂O emissions from wastewater effluent discharged to aquatic environments (Gg)
- 16 US_{POP} = U.S. population
- 17 US_{POPND} = U.S. population that is served by biological denitrification (from CWNS)
- 18 WWTP = Fraction of population using WWTP (as opposed to septic systems)
- 19 EF₁ = Emission factor (3.2 g N₂O/person-year) – plant with no intentional denitrification
- 20 EF₂ = Emission factor (7 g N₂O/person-year) – plant with intentional denitrification
- 21 Protein = Annual per capita protein consumption (kg/person/year)
- 22 F_{NPR} = Fraction of N in protein, default = 0.16 (kg N/kg protein)
- 23 F_{NON-CON} = Factor for non-consumed protein added to wastewater (1.4)
- 24 F_{IND-COM} = Factor for industrial and commercial co-discharged protein into the sewer system
- 25 (1.25)
- 26 N_{SLUDGE} = N removed with sludge, kg N/yr
- 27 EF₃ = Emission factor (0.005 kg N₂O -N/kg sewage-N produced) – from effluent
- 28 0.9 = Amount of nitrogen removed by denitrification systems (EPA 2008)
- 29 44/28 = Molecular weight ratio of N₂O to N₂

30 U.S. population data were taken from the U.S. Census Bureau International Database (U.S. Census 2012) and
 31 include the populations of the United States, American Samoa, Guam, Northern Mariana Islands, Puerto Rico, and
 32 the Virgin Islands. The fraction of the U.S. population using wastewater treatment plants is based on data from the
 33 1989, 1991, 1993, 1995, 1997, 1999, 2001, 2003, 2005, 2007, 2009, and 2011 American Housing Survey (U.S.
 34 Census 2011). Data for intervening years were obtained by linear interpolation. The emission factor (EF₁) used to
 35 estimate emissions from wastewater treatment for plants without intentional denitrification was taken from IPCC
 36 (2006), while the emission factor (EF₂) used to estimate emissions from wastewater treatment for plants with
 37 intentional denitrification was taken from Scheehle and Doorn (2001). Data on annual per capita protein intake were
 38 provided by U.S. Department of Agriculture Economic Research Service (USDA 2009). Protein consumption data
 39 for 2005 through 2011 were extrapolated from data for 1990 through 2004. Table 8-15 presents the data for U.S.
 40 population and average protein intake. An emission factor to estimate emissions from effluent (EF₃) has not been
 41 specifically estimated for the United States, thus the default IPCC value (0.005 kg N₂O-N/kg sewage-N produced)
 42 was applied. The fraction of N in protein (0.16 kg N/kg protein) was also obtained from IPCC (2006). The factor
 43 for non-consumed protein and the factor for industrial and commercial co-discharged protein were obtained from
 44 IPCC (2006). Sludge generation was obtained from EPA (1999) for 1988, 1996, and 1998 and from Beecher et al.
 45 (2007) for 2004. Intervening years were interpolated, and estimates for 2005 through 2011 were forecasted from the
 46 rest of the time series. An estimate for the N removed as sludge (N_{SLUDGE}) was obtained by determining the amount
 47 of sludge disposed by incineration, by land application (agriculture or other), through surface disposal, in landfills,
 48 or through ocean dumping. In 2011, 277 Gg N was removed with sludge.

49 Table 8-15: U.S. Population (Millions), Available Protein (kg/person-year), and Protein Consumed (kg/person-year)

Year	Population	Available Protein	Protein Consumed
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1990	253	38.7		29.6
2005	300	41.7		32.0
2007	305	42.1		32.3
2008	308	42.2		32.4
2009	311	42.4		32.5
2010	313	42.6		32.7
2011	316	42.8		32.8

Source: U.S. Census Bureau 2012, USDA 2009.

1 Table 8-16: Fate of Sludge Removed by Domestic Wastewater Treatment

Disposal Practices						
Distribution (1000 kg N)	1990	1995	2000	2005	2010	2011
Incineration	35,027.35	37,806.16	38,399.04	38,595.85	38,301.05	38,215.54
Land Application	77,378.34	97,230.98	113,311.73	129,196.74	144,113.04	147,054.99
<i>Ag</i>	52,198.15	69,001.16	83,522.63	98,080.96	112,014.99	114,778.24
<i>Other</i>	25,180.19	28,229.81	29,789.11	31,115.78	32,098.05	32,276.75
Surface Disposal	20,325.19	16,142.13	10,243.93	4,586.01	2,558.71	2,275.43
Landfill	72,962.21	75,945.15	74,158.54	71,407.98	67,609.40	66,790.83
Ocean Dumping	8,294.65	-	-	-	-	-
Other	1,645.76	6,353.98	11,312.32	16,478.76	21,661.26	22,702.30

2 Uncertainty and Time-Series Consistency

3 The overall uncertainty associated with both the 2011 CH₄ and N₂O emission estimates from wastewater treatment
4 and discharge was calculated using the IPCC Good Practice Guidance Tier 2 methodology (2000). Uncertainty
5 associated with the parameters used to estimate CH₄ emissions include that of numerous input variables used to
6 model emissions from domestic wastewater, and wastewater from pulp and paper manufacture, meat and poultry
7 processing, fruits and vegetable processing, ethanol production, and petroleum refining. Uncertainty associated with
8 the parameters used to estimate N₂O emissions include that of sewage sludge disposal, total U.S. population,
9 average protein consumed per person, fraction of N in protein, non-consumption nitrogen factor, emission factors
10 per capita and per mass of sewage-N, and for the percentage of total population using centralized wastewater
11 treatment plants.

12 The results of this Tier 2 quantitative uncertainty analysis are summarized in Table 8-17. Methane emissions from
13 wastewater treatment were estimated to be between 11.5 and 20.7 Tg CO₂ Eq. at the 95 percent confidence level (or
14 in 19 out of 20 Monte Carlo Stochastic Simulations). This indicates a range of approximately 29 percent below to
15 28 percent above the 2011 emissions estimate of 16.2 Tg CO₂ Eq. N₂O emissions from wastewater treatment were
16 estimated to be between 1.2 and 10.2 Tg CO₂ Eq., which indicates a range of approximately 77 percent below to 97
17 percent above the 2011 emissions estimate of 5.2 Tg CO₂ Eq.

18 Table 8-17: Tier 2 Quantitative Uncertainty Estimates for CH₄ Emissions from Wastewater Treatment (Tg CO₂ Eq.
19 and Percent)

Source	Gas	2011 Emission Estimate (Tg CO₂ Eq.)	Uncertainty Range Relative to Emission Estimate^a			
			(Tg CO₂ Eq.)		(%)	
			Lower Bound	Upper Bound	Lower Bound	Upper Bound

Wastewater Treatment	CH₄	16.2	11.5	20.7	-29%	+28%
Domestic	CH ₄	7.6	5.6	9.6	-26%	+27%
Industrial	CH ₄	8.6	4.6	12.7	-47%	+48%
Wastewater Treatment	N₂O	5.2	1.2	10.2	-77%	+97%

^a Range of emission estimates predicted by Monte Carlo Stochastic Simulation for a 95 percent confidence interval.

1 Methodological recalculations were applied to the entire time-series to ensure time-series consistency from 1990
2 through 2011. Details on the emission trends through time are described in more detail in the Methodology section,
3 above.

4 QA/QC and Verification

5 A QA/QC analysis was performed on activity data, documentation, and emission calculations. This effort included a
6 Tier 1 analysis, including the following checks:

- 7 • Checked for transcription errors in data input;
- 8 • Ensured references were specified for all activity data used in the calculations;
- 9 • Checked a sample of each emission calculation used for the source category;
- 10 • Checked that parameter and emission units were correctly recorded and that appropriate conversion factors
11 were used;
- 12 • Checked for temporal consistency in time series input data for each portion of the source category;
- 13 • Confirmed that estimates were calculated and reported for all portions of the source category and for all years;
- 14 • Investigated data gaps that affected emissions estimates trends; and
- 15 • Compared estimates to previous estimates to identify significant changes.

16 All transcription errors identified were corrected. The QA/QC analysis did not reveal any systemic inaccuracies or
17 incorrect input values.

18 Recalculations Discussion

19 Production data were updated to reflect updated USDA NASS datasets. This resulted in minor changes to the
20 emission estimates from the previous inventory. In addition, population updates from the U.S. Census resulted in
21 minor changes to domestic wastewater treatment emission estimates from 2000 through 2010.

22 Planned Improvements

23 The methodology to estimate CH₄ emissions from domestic wastewater treatment currently utilizes estimates for the
24 percentage of centrally treated wastewater that is treated by aerobic systems and anaerobic systems. These data
25 come from the 1992, 1996, 2000, and 2004 CWNS. The question of whether activity data for wastewater treatment
26 systems are sufficient across the time series to further differentiate aerobic systems with the potential to generate
27 small amounts of CH₄ (aerobic lagoons) versus other types of aerobic systems, and to differentiate between
28 anaerobic systems to allow for the use of different MCFs for different types of anaerobic treatment systems,
29 continues to be explored. The CWNS data for 2008 were evaluated for incorporation into the inventory, but due to
30 significant changes in format, this dataset is not sufficiently detailed for inventory calculations. However, additional
31 information and other data continue to be evaluated to update future years of the Inventory.

32 For industrial wastewater emissions, data recently collected by EPA's Office of Air for pulp and paper mills and
33 petroleum refineries is being evaluated to determine if sufficient information is available to update the estimates of
34 wastewater generated per unit of production and the percent of industry wastewater treated anaerobically in these
35 industries (%TA). Initial evaluations of EPA's Office of Air data for pulp and paper manufacturing indicate there is
36 sufficient information to update emission estimates in the next inventory year. Data collected in 2012 under the
37 EPA's GHGRP will also be investigated for updating this variable. In examining data from EPA's GHGRP for use
38 in improving the emission estimates for the industrial wastewater category, particular attention will be made to
39 ensure time series consistency, as the facility-level reporting data from EPA's GHGRP are not available for all
40 inventory years as reported in this inventory. In implementing improvements and integration of data from EPA's
41 GHGRP, the latest guidance from the IPCC on the use of facility-level data in national inventories will be relied

1 upon.²⁵² For all industries, EPA will continue to review new research on industrial wastewater characteristics,
2 utilization of treatment systems, and associated greenhouse gas emissions as it becomes available. Before the
3 incorporation of any new data, EPA will ensure it is representative of industry conditions.

4 Currently, it is assumed that all aerobic wastewater treatment systems are well managed and produce no CH₄ and
5 that all anaerobic systems have an MCF of 0.8. Efforts to obtain better data reflecting emissions from various types
6 of municipal treatment systems are currently being pursued.

7 With respect to estimating N₂O emissions, the default emission factors for indirect N₂O from wastewater effluent
8 and direct N₂O from centralized wastewater treatment facilities have a high uncertainty. Research is being
9 conducted by WERF to measure N₂O emissions from municipal treatment systems. In addition, a literature review
10 has been conducted focused on N₂O emissions from wastewater treatment to determine the state of such research
11 and identify data to develop a country-specific N₂O emission factor or alternate emission factor or method. Such
12 data will continue to be reviewed as they are available to determine if a country-specific N₂O emission factor can or
13 should be developed, or if alternate emission factors should be used.

14 Previously, EPA used new measurement data from WERF to develop U.S.-specific emission factors for CH₄
15 emissions from septic systems and incorporated it into the inventory emissions calculation. Due to the high
16 uncertainty of the measurements for N₂O from septic systems, estimates of N₂O emissions were not included.
17 Appropriate emission factors for septic system N₂O emissions will continue to be investigated as the data collected
18 by WERF indicate that septic soil systems are a source of N₂O emissions.

19 In addition, the estimate of N entering municipal treatment systems is under review. The factor that accounts for
20 non-sewage N in wastewater (bath, laundry, kitchen, industrial components) also has a high uncertainty. Obtaining
21 data on the changes in average influent N concentrations to centralized treatment systems over the time series would
22 improve the estimate of total N entering the system, which would reduce or eliminate the need for other factors for
23 non-consumed protein or industrial flow. The dataset previously provided by the National Association of Clean
24 Water Agencies (NACWA) was reviewed to determine if it was representative of the larger population of
25 centralized treatment plants for potential inclusion into the inventory. However, this limited dataset was not
26 representative of the number of systems by state or the service populations served in the United States, and therefore
27 could not be incorporated into the inventory methodology. Additional data sources will continue to be researched
28 with the goal of improving the uncertainty of the estimate of N entering municipal treatment systems.

29 The value used for N content of sludge continues to be investigated. This value is driving the N₂O emissions for
30 wastewater treatment and is static over the time series. To date, new data has not been identified that would be able
31 to establish a time series for this value. The amount of sludge produced and sludge disposal practices will also be
32 investigated. In addition, based on UNFCCC review comments, improving the transparency of the fate of sludge
33 produced in wastewater treatment will also be investigated.

34 A review of other industrial wastewater treatment sources for those industries believed to discharge significant loads
35 of BOD and COD has been ongoing. Food processing industries have the highest potential for CH₄ generation due
36 to the waste characteristics generated, and the greater likelihood to treat the wastes anaerobically. However, in all
37 cases there is dated information available on U.S. treatment operations for these industries. Previously, organic
38 chemicals, the seafood processing industry and coffee processing were investigated to estimate their potential to
39 generate CH₄. Due to the insignificant amount of CH₄ estimated to be emitted and the lack of reliable, up-to-date
40 data, these industries were not selected for inclusion in the industry. Preliminary analyses of the beer and malt and
41 dairy products industries has been performed. These industries will continue to be investigated for incorporation.
42 Other industries will be reviewed as necessary for inclusion in future years of the Inventory using EPA's Permit
43 Compliance System and Toxics Release inventory.

44 In addition, available datasets will be reviewed to provide further information on the fates of sludge removed by
45 domestic wastewater treatment in the next Inventory report.

²⁵² See: http://www.ipcc-nggip.iges.or.jp/meeting/pdfiles/1008_Model_and_Facility_Level_Data_Report.pdf.

8.3. Composting (IPCC Source Category 6D)

Composting of organic waste, such as food waste, garden (yard) and park waste, and sludge, is common in the United States. Advantages of composting include reduced volume in the waste material, stabilization of the waste, and destruction of pathogens in the waste material. The end products of composting, depending on its quality, can be recycled as fertilizer and soil amendment, or be disposed in a landfill. Approximately 400 composting facilities operate in the United States (WBJ 2010).

Composting is an aerobic process and a large fraction of the degradable organic carbon in the waste material is converted into carbon dioxide (CO₂). Methane (CH₄) is formed in anaerobic sections of the compost, but it is oxidized to a large extent in the aerobic sections of the compost. Anaerobic sections are created in composting piles when there is excessive moisture or inadequate aeration (or mixing) of the compost pile. The estimated CH₄ 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 the N content of the feedstock and how well the compost pile is managed, nitrous oxide (N₂O) emissions can be produced. The formation of N₂O is complicated, but is mainly associated with anaerobic conditions. Emissions vary and range from less than 0.5 percent to 5 percent of the initial content of the material (IPCC 2006).

From 1990 to 2011, the amount of material composted in the United States has increased from 3,810 Gg to 18,449 Gg, an increase of approximately 384 percent. From 2000 to 2011, the amount of material composted in the United States has increased by approximately 24 percent. Emissions of CH₄ and N₂O from composting have increased by the same percentage. In 2011, CH₄ emissions from composting (see Table 8-18 and Table 8-19) were 1.5 Tg CO₂ Eq. (74 Gg), and N₂O emissions from composting were 1.7 Tg CO₂ Eq. (5.5 Gg). The wastes composted primarily include yard trimmings (grass, leaves, and tree and brush trimmings) and food scraps from residences and commercial establishments (such as grocery stores, restaurants, and school and factory cafeterias). The composted waste quantities reported here do not include backyard composting. The growth in composting since the 1990s is attributable to primarily two factors: (1) steady growth in population and residential housing, and (2) the enactment of legislation by state and local governments that discouraged the disposal of yard trimmings in landfills. In 1992, 11 states and the District of Columbia had legislation in effect that banned or discouraged disposal of yard trimmings in landfills. Currently, 23 states and the District of Columbia, representing about 50 percent of the nation's population, have enacted such legislation (EPA 2010). The total amount of waste composted has decreased slightly since 2008, by approximately 8 percent.

Table 8-18: CH₄ and N₂O Emissions from Composting (Tg CO₂ Eq.)

Activity	1990	2005	2007	2008	2009	2010	2011
CH ₄	0.3	1.6	1.7	1.7	1.6	1.5	1.5
N ₂ O	0.4	1.7	1.8	1.9	1.8	1.7	1.7
Total	0.7	3.3	3.5	3.5	3.3	3.2	3.3

Table 8-19: CH₄ and N₂O Emissions from Composting (Gg)

Activity	1990	2005	2007	2008	2009	2010	2011
CH ₄	15	75	79	80	75	73	74
N ₂ O	1	6	6	6	6	5	6

Methodology

Methane and N₂O emissions from composting depend on factors such as the type of waste composted, the amount and type of supporting material (such as wood chips and peat) used, temperature, moisture content and aeration during the process.

The emissions shown in Table 8-18 and Table 8-19 were estimated using the IPCC default (Tier 1) methodology (IPCC 2006), which is the product of an emission factor and the mass of organic waste composted (note: no CH₄

1 recovery is expected to occur at composting operations):

$$2 \quad E_i = M \times EF_i$$

3 where,

- 4 E_i = CH₄ or N₂O emissions from composting, Gg CH₄ or N₂O,
- 5 M = mass of organic waste composted in Gg,
- 6 EF_i = emission factor for composting, 4 g CH₄/kg of waste treated (wet basis) and 0.3 g
- 7 N₂O/kg of waste treated (wet basis) (IPCC 2006), and
- 8 i = designates either CH₄ or N₂O.

9 Estimates of the quantity of waste composted (M) are presented in Table 8-20. Estimates of the quantity composted
 10 for 1990 and 1995 were taken from the *Characterization of Municipal Solid Waste in the United States: 1996*
 11 *Update* (Franklin Associates 1997); estimates of the quantity composted for 2000, 2005, 2006, 2007, 2008, and 2009
 12 were taken from EPA’s *Municipal Solid Waste In The United States: 2009 Facts and Figures* (EPA 2010);
 13 estimates of the quantity composted for 2010 were taken from EPA’s *Municipal Solid Waste In The United States:*
 14 *2010 Facts and Figures* (EPA 2011); estimates of the quantity composted for 2011 were calculated using the 2010
 15 quantity composted and a ratio of the U.S. population in 2010 and 2011 (U.S. Census Bureau 2012). The estimated
 16 quantity of waste composted in 2010 was revised based on updated information (EPA 2011).

17 Table 8-20: U.S. Waste Composted (Gg)

Activity	1990	2005	2007	2008	2009	2010	2011
Waste							
Composted	3,810	18,643	19,695	20,049	18,824	18,298	18,449

Source: EPA 2008 and EPA 2011.

18 Uncertainty and Time-Series Consistency

19 Little is known about the site-specific operating conditions at the composting facilities in the United States. The
 20 generation of CH₄ and N₂O emissions is highly dependent on the characteristics of the feedstock material (e.g.,
 21 moisture content, C to N ratio, size), on the climate, and on the operating and maintenance practices (e.g., use of a
 22 shredder/grinder to maintain consistency in size of the feedstock material, frequency of pile rotation, addition of
 23 moisture, application of finished compost on the pile). The estimated uncertainty from the 2006 IPCC Guidelines is
 24 ±50 percent for the Tier 1 methodology. Emissions from composting in 2011 were estimated to be between 1.6 and
 25 4.9 Tg CO₂ Eq., which indicates a range of 50 percent below to 50 percent above the actual 2011 emission estimate
 26 of 3.3 Tg CO₂ Eq. (see Table 8-21).

27 Table 8-21 : Tier 1 Quantitative Uncertainty Estimates for Emissions from Composting (Tg CO₂ Eq. and Percent)

Source	Gas	2011 Emission Estimate (Tg CO ₂ Eq.)	Uncertainty Range Relative to Emission Estimate			
			(Tg CO ₂ Eq.)		(%)	
			Lower Bound	Upper Bound	Lower Bound	Upper Bound
Composting	CH ₄ , N ₂ O	3.3	1.6	4.9	-50%	+50%

28 Methodological recalculations were applied to the entire time series to ensure time-series consistency from 1990
 29 through 2011. Details on the emission trends through time are described in more detail in the Methodology section,
 30 above.

31 QA/QC and Verification

32 A QA/QC analysis was performed for data gathering and input, documentation, and calculation. A primary focus of
 33 the QA/QC checks was to ensure that the amount of waste composted annually was correct according to the latest
 34 EPA *Municipal Solid Waste In The United States: Facts and Figures* report.

1 **Recalculations Discussion**

2 The estimated amount of waste composted in 2010 was updated based on new data contained in EPA’s *Municipal*
3 *Solid Waste In The United States: 2010 Facts and Figures* (EPA 2011). The amounts of CH₄ and N₂O emissions
4 estimates presented in Table 8-18 and Table 8-19 were revised accordingly.

5 **Planned Improvements**

6 For future Inventories, additional efforts will be made to improve the estimates of CH₄ and N₂O emissions from
7 composting. For example, a literature search may be conducted to determine if emission factors specific to various
8 composting systems and composted materials are available. Further cooperation with estimating emissions in
9 cooperation with the LULUCF Other section will be made.

10 **8.4. Waste Sources of Indirect Greenhouse Gases**

11 In addition to the main greenhouse gases addressed above, waste generating and handling processes are also sources
12 of indirect greenhouse gas emissions. Total emissions of NO_x, CO, and NMVOCs from waste sources for the years
13 1990 through 2011 are provided in Table 8-22.

14 Table 8-22: Emissions of NO_x, CO, and NMVOC from Waste (Gg)

Gas/Source	1990	2005	2007	2008	2009	2010	2011
NO_x	+	2	2	2	2	2	2
Landfills	+	2	2	2	2	2	2
Wastewater Treatment	+	+	+	+	+	+	+
Miscellaneous ^a	+	+	+	+	+	+	+
CO	1	7	7	7	7	7	7
Landfills	1	6	6	6	6	6	6
Wastewater Treatment	+	+	+	+	+	+	+
Miscellaneous ^a	+	+	+	+	+	+	+
NMVOCs	673	114	111	109	76	76	76
Wastewater Treatment	57	49	48	47	33	33	33
Miscellaneous ^a	557	43	42	41	29	29	29
Landfills	58	22	21	21	14	14	14

^a 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.

+ Does not exceed 0.5 Gg.

15 **Methodology**

16 Due to the lack of data available at the time of publication, emission estimates for 2010 and 2011 rely on 2009 data
17 as a proxy. Emission estimates for 2009 were obtained from preliminary data (EPA 2010, EPA 2009), and
18 disaggregated based on EPA (2003), which, in its final iteration, will be published on the National Emission
19 Inventory (NEI) Air Pollutant Emission Trends web site. Emission estimates of these gases were provided by
20 sector, using a “top down” estimating procedure—emissions were calculated either for individual sources or for
21 many sources combined, using basic activity data (e.g., the amount of raw material processed) as an indicator of
22 emissions. National activity data were collected for individual source categories from various agencies. Depending
23 on the source category, these basic activity data may include data on production, fuel deliveries, raw material
24 processed, etc.

25 **Uncertainty and Time-Series Consistency**

26 No quantitative estimates of uncertainty were calculated for this source category. Methodological recalculations
27 were applied to the entire time-series to ensure time-series consistency from 1990 through 2011.

28

29