Better Understanding Wastewater Treatment’s Nitrous Oxide Emissions

The case for revised reporting protocols using variable emission factors based on nitrogen removal

INTRODUCTION
Nitrous oxide (N\textsubscript{2}O) is a major contributor to direct (Scope 1) greenhouse gas (GHG) emissions from wastewater treatment plants (WWTPs). Activated sludge systems typically represent the bulk of mainstream treatment processes for municipal WWTPs, both in Australia and most developed countries. N\textsubscript{2}O has been measured in the off-gases from such plants in many different locations. The underlying biochemical mechanisms are complex. The measured actual emission rates show high variability in N\textsubscript{2}O emissions, both in time and space (within a given bioreactor system) and between different bioreactors across a range of WWTP configurations. Under current GHG reporting protocols, both internationally and in Australia, fixed emission factors for N\textsubscript{2}O are typically used (i.e., invariant over time and source location). However, in the calculations underlying the emissions reported under different protocols, the way in which these factors are defined and applied can be different. Hence, aside from the complexity and variability of the actual biochemical mechanisms at play, emission factor definition and calculation methodology are a first potential source of confusion when reporting N\textsubscript{2}O emissions from WWTPs. A second source of potential confusion stems from the use of fixed emission factors (i.e., without reference to type of treatment plant or its performance, for example, in terms of nitrogen removal). A third source of confusion concerns the boundary of the WWTP and the distinction between N\textsubscript{2}O emissions from the WWTP treatment processes per se versus those associated with the discharge of WWTP effluent to a receiving water body (e.g., river, estuary ocean) or land (in the case of irrigation), and the disposal of biosolids.

This paper focusses on the first two potential sources of confusion around WWTP N\textsubscript{2}O emissions mentioned above. It starts with an overview, in simple terms, of the main biochemical mechanisms of N\textsubscript{2}O formation in activated sludge systems, based on current knowledge from research literature. Next, recent results (from both published literature and previously unpublished data, both overseas and in Australia) are examined, highlighting a common trend in average N\textsubscript{2}O emission factors from different WWTPs. Lastly, these results are compared with the N\textsubscript{2}O emission factors as defined in the IPCC (2019) and the current Australian NGER Determination (2020) reporting protocols.

Keywords: Nitrous oxide, N\textsubscript{2}O, greenhouse gas emissions reporting, emission factors, wastewater treatment plants

LITERATURE REVIEW
N\textsubscript{2}O formation biochemical mechanisms in wastewater treatment
N\textsubscript{2}O formation in WWTPs have been previously reviewed by others (inter alia Law et al., 2012; Wunderlin et al., 2012, 2013; Valkova et al., 2021). In their review, Law et al. (2012) made following general observations:

- The microbial nitrogen transformation processes in a WWTP are fundamentally the same as in other environments such as soil, marine and freshwater
habitats. However, unlike most other environments, wastewater treatment plants are engineered systems designed to achieve high nitrogen conversion rates.

- Domestic wastewater usually contains relatively high concentrations of nitrogen (N), around 20–70 mg/L total nitrogen (TN) as N. To achieve almost complete nitrogen removal within short retention times (in the order of hours), high reactor N loading rates are applied, resulting in relatively high nitrification and denitrification rates, and potentially relatively high rates of N₂O production.
- Bacterial communities in a WWTP are subjected to rapid changes in process conditions that are applied to promote aerobic or anoxic biochemical reactions. Such rapid changes likely cause physiological stress on both the nitrifying and denitrifying communities, and the potential to induce transient behaviours like N₂O formation.
- Active aeration is used to induce aerobic conditions in WWTPs. The aeration systems are engineered to efficiently transfer oxygen to the bioreactor, but also enable efficient transfer of N₂O from the liquid phase to the gas phase. Therefore, any temporary imbalance between N₂O production and consumption in the bioreactor could result in N₂O accumulation and/or stripping during aeration.
- Given that WWTPs are highly engineered, there are opportunities to mitigate N₂O emissions by improving process design and/or operation.

The main pathways for N₂O formation, based on the research literature of underlying biochemical mechanisms are summarised in Table 1. The two main pathways are autotrophic nitrification (via nitrifier denitrification; and/or hydroxylamine oxidation) and heterotrophic denitrification, with a third (chemical) pathway considered to be uncommon.

Table 1: Summary of main biochemical pathways for nitrous oxide formation in wastewater treatment, typified by activated sludge systems.

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Sub-pathway</th>
<th>Process conditions favouring N₂O formation</th>
<th>Microorganisms involved</th>
<th>Underlying biochemical mechanism (key points)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Autotrophic nitrification</td>
<td>a. Nitrifier denitrification</td>
<td>Low dissolved oxygen (DO) concentrations and accumulation of nitrite</td>
<td>Mainly ammonia oxidising bacteria (AOB), i.e., nitrifiers oxidising ammonia (NH₄⁺/NH₃) to nitrite (NO₂⁻)</td>
<td>AOB oxidise ammonia to nitrite (NO₃⁻), via hydroxylamine, and survive stress conditions (low DO, increased NO₂⁻ concentrations) by reducing NO₂⁻ to NO → N₂O, but lack N₂O reductase enzyme (to N₂), leading to N₂O production and emission.</td>
</tr>
<tr>
<td></td>
<td>b. Hydroxylamine oxidation</td>
<td>High DO, high turnover rates of AOB (high ammonium loading rates) and accumulation of hydroxylamine</td>
<td>Hydroxylamine is formed as an intermediate in the pathway for ammonia oxidation to nitrite (NO₂⁻) (see above). The mechanism for oxidation of hydroxylamine to NO₂⁻ includes formation of an unstable intermediate (nitrosyl radical, NOH). N₂O appears to be produced and emitted due to the unstable breakdown of NOH.</td>
<td></td>
</tr>
<tr>
<td>II. Heterotrophic denitrification</td>
<td></td>
<td>Anoxic conditions (low or zero DO), low pH, presence of hydrogen sulphide; accumulation of nitric oxide (NO) and nitrite (NO₂⁻); lack of biodegradable organic carbon (high TKN/COD ratio)</td>
<td>Heterotrophic denitrifiers (Partial) inhibition of N₂O reductase (enzyme).</td>
<td>N₂O reduction is an obligate final step in the four-step reduction chain for denitrification of nitrate (NO₃⁻ → NO₂⁻ → NO → N₂O → N₂ (gas)) Note 1</td>
</tr>
<tr>
<td>Chemical (uncommon)</td>
<td></td>
<td>High hydroxylamine concentrations and nitrite (typically in side-stream processes), very low pH</td>
<td>None/ Abiotic</td>
<td>Chemical</td>
</tr>
</tbody>
</table>

Note 1: There is some evidence (at least at laboratory scale) that heterotrophic denitrifiers have the capacity to act not only a source but also a sink for N₂O (i.e., to produce net N₂O under some conditions, due instability in the reduction chain, but net removal of N₂O under other conditions through denitrification of N₂O to N₂ gas).

Sources: Goreau et al. (1980); Igarashi et al. (1997); Law et al. (2012); Ribera-Guardia et al. (2014); Solen-Jofra et al. (2016); Stein et al. (2011); Tallec et al. (2006); Valkova et al. (2021); von Schulthess et al. (1995); Wunderlin et al. (2013).
Two GHG reporting protocols for domestic wastewater treatment were reviewed, namely: the 2019 Refinement to the 2006 IPCC Guidelines for National Greenhouse Gas Inventories (international); and the National Greenhouse and Energy Reporting (Measurement) Determination 2008 (2020 compilation), which is applicable in Australia (Australian Govt, 2020).

In respect of N₂O for domestic wastewater treated in centralised ‘aerobic’ treatment plants (not including effluent discharge), the IPCC guidelines (IPCC, 2019) apply a plant-wide fixed (default) emission factor (EFₙ₂ₒ⁻ₓₓₚₜ) of 0.016 kg N₂O-N/kg N (1.6%) with respect to the influent (raw wastewater) total nitrogen (TN) load - N₂O emissions are noted as ‘variable’ with a wide possible EF range given (0.016% - 4.5%) (IPCC, 2019). The IPCC default EFₙ₂ₒ⁻ₓₓₚₜ does not change with TN removal across the WWTP.

The origin of the IPCC (2019) default emission factor (EFₙ₂ₒ⁻ₓₓₚₜ = 0.016 kg N₂O-N/kg N) is presented in Figure 1, found in Annex 6A.5 of the IPCC 2019 Refinement guidelines. The data for this chart was sourced from various literature references (cited by IPCC, 2019) for 30 WWTPs considered to be reflective of ‘the most typical and widely used treatment processes globally’ (IPCC, 2019). It is a reasonable supposition that the average N₂O daily emission rate will be directly proportional to the TN mass loading rate (i.e., nominally the size of the WWTP), as suggested by the linear trend in Figure 1. However, the calculated slope of the trendline in Figure 1 (from which the IPCC default emission factor stems) was likely influenced heavily by the data from less than ten plants (a third of the dataset) with TN loading rates >4,000 kgN/d (approximately) i.e., relatively large WWTPs treating population equivalents of >300,000, indicatively. For many settings, including most of regional Australia and New Zealand, such large plants would be uncommon in terms of number. Moreover, the inherent assumption behind the trendline in Figure 1 is that process performance across the selected WWTPs, at least in terms of N removal, is essentially the same. In reality, N removal performance can differ widely between different WWTPs. The underlying biochemical mechanisms (see Table 1) suggest that N₂O emissions are linked with N removal (i.e., nitrification-denitrification biological reactions) over a range of plant sizes from laboratory scale to full scale.

![Figure 1: Correlation between influent Total nitrogen (TN) and N₂O emissions (reproduced from IPCC (2019), Annex 6A.5)](image-url)
By contrast, for $N_2O$ from domestic wastewater treatment, the NGER Determination (Australian Govt., 2020) applies a plant-wide fixed EF$_{N2O,WWTP}$ of 2.082 tonnes CO$_2$-e as $N_2O$ per tonne N “produced” (removed), which converts to 0.005 tonnes N2O-N per tonne N removed (0.5% of TN removed) at 265 tonnes CO$_2$-e/tonne N$_2O$. For a case where treatment removal (by mass balance calculations in the NGER methodology) is indicatively 80% of influent TN, the NGER EF for $N_2O$ is equivalent to 0.4% of influent TN (i.e., four times lower than the IPCC default EF, see above). The NGER methodology provides no guidance on the range in the EF for $N_2O$. However, when comparing the IPCC and NGER methods in equivalent terms (i.e., both relative to influent TN), the NGER EF effectively decreases with decreasing TN removal across the WWTP.

In both protocols (IPCC and NGER), the remainder of influent TN not removed (i.e., effluent TN discharged) is separately accounted for using a different EF for discharge defined in each of the respective guidelines, to account for $N_2O$ expected to be produced and released from the receiving environment. This aspect was not studied here.

**RESULTS AND DISCUSSION**

Figure 2 shows that both the EU and AU datasets suggest linear trends in plant-wide EF$_{N2O,WWTP}$ where the EF is inversely related to TN removal (expressed in percentage terms) across the WWTP. The linear regression correlation is stronger for the EU dataset ($R^2 = 0.86$), which included more datapoints ($n = 20$), compared with the AU dataset ($R^2 = 0.56$, $n = 7$). One AU WWTP was excluded from the regression analysis since this measurement campaign was a special case (see Figure 2). The measured emissions were relatively low for this plant (considering its configuration and extent of TN removal, compared with other similar plants). Possible reasons for this were related to methodological differences around the off-gas measurement, and the presence of a thick scum/foam layer on the surface of the aeration tanks in the vicinity of the gas hoods for off-gas collection. Biological removal of $N_2O$ through heterotrophic denitrification in the scum/foam layer is possible.

The trendlines identified in Figure 2 imply that, in relative terms, $N_2O$ emissions increase as N removal decreases (i.e., where nitrification-denitrification reactions are incomplete or partial). Conversely, lower $N_2O$ emissions in relative terms, are expected to occur in WWTPs that achieve higher degrees of N removal (i.e., where nitrification-denitrification reactions are largely complete by the time the effluent leaves the bioreactors). Obviously, this trend has the inherent assumption that N removal occurs predominantly in one bioreactor (or one set of similar bioreactors). In practice, within the same WWTP, it is possible that different bioreactors (representing sequential or parallel process steps) emit $N_2O$ emissions to different degrees and remove N to different degrees, whereas the WWTP overall achieves a relatively high percentage of TN removal. In such cases, the WWTP data might give excursions from the trend in Figure 2.

As explained by Valkova et al. (2021), there are fundamental reasons why $N_2O$ emissions might be linked to the extent of TN removal across the mainstream processes for WWTPs. Essentially, most of the emitted $N_2O$ is attributed to autotrophic nitrification pathways. The biochemical mechanisms of $N_2O$ formation during nitrification are associated with higher ammonia oxidation rates, and/or higher reactor prevalence of partial nitrification/denitrification products (e.g., nitrite), both of which are linked with higher reactor nitrogen loading rates, spatial feed...
distribution, DO concentrations, and transient aerobic conditions (e.g., at the start of aeration, moving from low to higher reactor DO). In each case, these conditions typically tend to occur in plants designed or operated for lower degrees of TN removal and/or shorter sludge ages. Plants that achieve near-complete TN removal tend to be designed for longer sludge ages and have larger reactors that are more lightly loaded, with larger anoxic fractions ($N_2O$ sinks).

Given the limitations from variance and number of datapoints in the datasets, Figure 2 suggests that the average $EF_{N2O-WWTP}$, for a given degree of TN removal, is higher in the AU WWTPs (UQ dataset) than the EU WWTPs (Valkova et al., 2021 dataset). Many factors might contribute to variability in $N_2O$ emissions, for a given degree of TN removal (i.e., the vertical scatter of datapoints in Figure 2). Some of these might be physical factors around plant design and operation (e.g., aeration and recycles rates, reactor dissolved oxygen concentration, reactor type and configuration including compartmentalisation, feed and recycle set-up, flow and/or diurnal load patterns, reactor temperature etc.), while others might be biological factors (e.g., wastewater composition, microbial community diversity). Furthermore, given the difficulty of measuring off-gas or dissolved $N_2O$ in full scale reactors, methodological factors (systematic or measurement errors) likely also contribute to the EF uncertainty.

To compare reporting methodologies, the $N_2O$ EF$_{WWTP}$ can be expressed in two ways: (1) relative to influent TN load (e.g., IPCC, 2019); or (2) relative to TN removed (e.g., NGER, Australian Govt., 2020). Both approaches were examined for the datasets considered in this study.

**$N_2O$ EF relative to Influent TN load**

Figure 3 relates the actual $N_2O$ emissions data (from Figure 2) to the default emissions factors derived from the IPCC and NGER (equivalent) reporting protocols, expressed on a common basis, namely, $N_2O$ emissions with respect to percent influent TN load vs. percent TN removal across the whole WWTP. It shows that the IPCC EF (1.6%) is very conservative (i.e., high), makes no provision for the degree of TN removal and likely over-estimates actual $N_2O$ emissions for typical WWTPs. Conversely, the NGER EF (0.5% relative to TN removed in the 2020 compilation; decreased from 1.0% in previous NGER compilations) might only be representative of $N_2O$ emissions for WWTPs achieving at least approximately 80% TN removal. The formulation of the NGER EF as constant relative to TN removed results in the equivalent EF (relative to influent TN) decreasing with decreasing degrees of TN removal, whereas the actual $N_2O$ data suggests the opposite trend. The NGER methodology will therefore likely underestimate actual $N_2O$ emissions for WWTPs that achieve lower degrees of TN removal (indicatively <83% at $EF_{N2O-WWTP} = 0.4\%$ in Figure 3).

Table 1 shows the outcome of correcting the EF’s applied in the IPCC and NGER methodologies, based on the ‘best fit’ linear regression for the pooled EU and AU datasets (excluding 1 no. AU WWTP, see above), as shown in Figure 4. The linear correlation for the pooled datasets is relatively weak ($r^2 = 0.55$), as expected, given that the EU and AU datasets appear to be distinct from each other (Figure 2). Further work is required to understand the underlying reasons for such dataset differences, be they process related (e.g., temperature, type and configuration of bioreactors) or methodological (e.g., around $N_2O$ measurement campaigns, and/or TN removal calculation for the bioreactors per se or the WWTP as a whole).

**$N_2O$ EF relative to TN removed**

Since the predominant mechanisms for $N_2O$ formation in WWTPs involve biological pathways for nitrification and denitrification (Table 1), it is more sensible to relate the $N_2O$ emission factor to TN removal than influent TN.

Figure 5 presents the same datasets as for Figure 2, except with the emission factor (y-axis) expressed as a percentage relative to TN removed. Figure 5 shows improved linear correlations for both the EU and AU datasets, compared with Figure 2.

Figure 6 plots the actual $N_2O$ emissions data (from Figure 5) to the default emissions factors derived from the IPCC (equivalent) and NGER reporting protocols, expressed on the alternative common basis, namely, $N_2O$ emissions as a percentage of TN removed vs. percent TN removal across the whole WWTP. In this case, the IPCC equivalent emission factor (which is constant with respect to influent TN load, see Figure 3) increases as TN removal decreases (Figure 6) and at 100% TN removal (theoretically) is numerically the same in percentage terms as the IPCC default (1.6% of influent TN load). The actual $N_2O$ data from both the EU and AU datasets reviewed here (Figure 6) suggest that the IPCC default is too conservative (too high).
The NGER default factor (constant with respect to TN removed, see Figure 6) might be more appropriate for WWTPs with TN removal in the higher range (indicatively 80 to 97%), but is likely to over-predict N\textsubscript{2}O emissions for some plants (e.g., most plants in the EU dataset in this range). Conversely, the data suggest that the NGER default factor is likely to under-predict N\textsubscript{2}O emissions in the lower range of TN removal (indicatively 60% to 80%), potentially by a large margin. The best approach would be to adopt a variable N\textsubscript{2}O emission factor, expressed with respect to percent TN removed (i.e., like the NGER approach) whereby the emission factor is a function of percent TN removal across the whole WWTP. Figure 7 shows a plot of the emission factor expressed on this basis for the pooled datasets (EU and AU) reviewed in this study. The correlation ($r^2 = 0.66$) is somewhat better than that in Figure 4, but with room for improvement, potentially by adding more data from further field measurement campaigns (in Australia and internationally) and standardising measurement methodologies.

Finally, for context, 2015-16 TN removal data from a benchmarking study of WWTPs in Australia and New Zealand (de Haas et al., 2018) is summarised in Table 3. There is a wide variation in the range of actual TN removal for the WWTPs surveyed in this benchmarking study. Adopting the simple average of TN removal (83%) of all the WWTP surveyed (in 2015-16) might suggest that the NGER default factor is appropriate. However, using a weighted average of 56%TN removal (from Table 3 weighted on plant size i.e., EP, and hence mass of TN treated), by extrapolation from Figure 7, at national scale the NGER default factor is potentially under-estimating N\textsubscript{2}O emissions by a factor of more than four-fold. This could be corrected by adopting a variable emission factor for N\textsubscript{2}O reporting, based on TN removed for each plant, as suggested above.

**CONCLUSION**

Actual (measured) emissions data from the mainstream processes (typically activated sludge) of WWTPs in Europe and Australia suggest that N\textsubscript{2}O emissions (expressed relative to influent TN load), on average, increase as the degree of TN removal across the plant decreases. That is, less N\textsubscript{2}O is likely to be emitted as overall TN removal improves. The average N\textsubscript{2}O emission factor is likely to increase by at least 2 to 5-fold (and up to 10-fold or more) as the degree of TN removal decreases from >90% to <70%. As suggested recently by Valkova et al. (2021), linking the N\textsubscript{2}O emission factor to the degree of TN removal across a WWTP, is recommended as a better approach for GHG accounting than the use of fixed emission factors in current reporting protocols (e.g., IPCC and NGER). Additional full-scale measurement campaigns for N\textsubscript{2}O emissions from WWTPs, along with alignment of measurement and reporting methodologies, are needed to confirm the trends highlighted in this study. A review of the IPCC and NGER reporting methodologies for wastewater treatment N\textsubscript{2}O emissions is recommended in the light of these findings. In light of the identified trend toward N\textsubscript{2}O emissions with improved N removal, future work should include re-examination of the trade-offs for WWTPs between improved N removal (through more advanced treatment) and total GHG emissions. The total GHG footprint will take into account both direct emissions (including N\textsubscript{2}O and methane, potentially) and indirect emissions (i.e. grid electricity use and chemicals use, biosolids transport/ disposal etc), noting local or regional differences around indirect emission factors (e.g. the renewable energy component of grid electricity, and biosolids handling).
Figure 2: Comparison of average nitrous oxide emission factor for wastewater treatment (EF$_{N2O-WWTP}$), defined as percent of influent TN load, versus percent of influent TN removal by the WWTP for European (EU) and Australian (AU) datasets. Note: One AU WWTP (Special case) excluded from regression analysis.
Figure 3: Comparison of average nitrous oxide emission factor for wastewater treatment (EF$_{\text{N2O-WWTP}}$) from actual (measured) data with default values from IPCC or NGER protocols, related on an equivalent basis with respect to WWTP influent TN.
Table 2: Summary of current and suggested corrected emission factors for N$_2$O from WWTPs

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>60</td>
<td>1.6</td>
<td>1.67</td>
<td>0.300</td>
<td>0.5</td>
<td>2.78</td>
</tr>
<tr>
<td>65</td>
<td>1.6</td>
<td>1.44</td>
<td>0.325</td>
<td>0.5</td>
<td>2.21</td>
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<tr>
<td>70</td>
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<td>1.21</td>
<td>0.350</td>
<td>0.5</td>
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<tr>
<td>75</td>
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<td>0.98</td>
<td>0.375</td>
<td>0.5</td>
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<tr>
<td>80</td>
<td>1.6</td>
<td>0.74</td>
<td>0.400</td>
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<td>0.93</td>
</tr>
<tr>
<td>85</td>
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<td>0.60</td>
</tr>
<tr>
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<td>0.28</td>
<td>0.450</td>
<td>0.5</td>
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<tr>
<td>95</td>
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<td>0.05</td>
<td>0.475</td>
<td>0.5</td>
<td>0.05</td>
</tr>
<tr>
<td>(100)*</td>
<td>1.6</td>
<td>0.00</td>
<td>0.500</td>
<td>0.5</td>
<td>0.00</td>
</tr>
</tbody>
</table>

* Hypothetical (Without advanced tertiary treatment, WWTP typically achieve up to approx. 97% TN removal)

Note 1: Based on the “best fit” linear regression of pooled actual datasets in Figure 4.

Note 2: Based on the “best fit” linear regression of pooled actual datasets in Figure 7.
Figure 4: Comparison of average nitrous oxide emission factor for wastewater treatment (EF$_{\text{N}_2\text{O-WWTP}}$) from actual (measured) data showing suggested corrected EF trendline from 'best fit' of pooled actual datasets, related on an equivalent basis of influent TN load.

\[ y = -0.0462x + 4.44 \\
R^2 = 0.5407 \]

- European WWTPs, Valkova et al. (2021)
- AU WWTPs (UQ data)
- IPCC (2019) default EF
- Corrected IPCC EF (All Data, indicative ‘best fit’)
Figure 5: Comparison of average nitrous oxide emission factor for wastewater treatment (EF\textsubscript{N2O-WWTP}), defined as percent of TN removed, versus percent of influent TN removal by the WWTP for European (EU) and Australian (AU) datasets. Note: One AU WWTP (Special case) excluded from regression analysis.
Figure 6: Comparison of average nitrous oxide emission factor for wastewater treatment (EF\textsubscript{N2O-WWTP}) from actual (measured) data with default values from IPCC or NGER protocols, related on an equivalent basis with respect to WWTP TN removed.
Figure 7: Comparison of average nitrous oxide emission factor for wastewater treatment ($\text{EF}_{\text{N}_2\text{O}-\text{WWTP}}$) from actual (measured) data showing suggested corrected EF trendline for pooled actual datasets, related on an equivalent basis with respect to TN removed.
Table 3: Summary of TN removal characteristics of WWTPs in Australia (including two in New Zealand) from 2015-16 benchmarking data

<table>
<thead>
<tr>
<th>WWTP Size Class (EP range)</th>
<th>Average of 2015-16 Adopted EP</th>
<th>No. of WWTPs (No. counted for TN removal)</th>
<th>Min. of %TN removal</th>
<th>Average of %TN removal</th>
<th>Max. of % TN removal</th>
<th>Wt Ave. %TN removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC1 (≤ 1,000)</td>
<td>482</td>
<td>22 (21)</td>
<td>65%</td>
<td>85%</td>
<td>97%</td>
<td>0.03%</td>
</tr>
<tr>
<td>SC2 (1,001 to 5,000)</td>
<td>2,414</td>
<td>47 (46)</td>
<td>46%</td>
<td>81%</td>
<td>98%</td>
<td>0.3%</td>
</tr>
<tr>
<td>SC3 (5,001 to 10,000)</td>
<td>7,591</td>
<td>28 (28)</td>
<td>52%</td>
<td>82%</td>
<td>98%</td>
<td>1%</td>
</tr>
<tr>
<td>SC4 (10,001 to 100,000)</td>
<td>38,267</td>
<td>107 (107)</td>
<td>19%</td>
<td>84%</td>
<td>98%</td>
<td>12%</td>
</tr>
<tr>
<td>SC5 (&gt; 100,000)</td>
<td>514,987</td>
<td>40 (37)</td>
<td>39%</td>
<td>86%</td>
<td>98%</td>
<td>43%</td>
</tr>
<tr>
<td>ALL</td>
<td>102,585</td>
<td>244 (239)</td>
<td>19%</td>
<td>83%</td>
<td>98%</td>
<td>56%</td>
</tr>
</tbody>
</table>

Excludes WWTPs for which non-valid (negative) TN removal data or <19% TN removal was reported

Note 1: Weighted average %TN removal contribution per Size Class (SC) to overall weighted average (ALL). Weighting from Adopted EP per WWTP relative to Total EP (for ALL plants).

EP: Equivalent population (persons) based on influent loading (COD or BOD and TKN)

Data sourced from de Haas et al. (2018)

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