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# Design of anaerobic membrane bioreactors for the valorization of dilute organic carbon waste streams<sup>†</sup>

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Aqueous waste organics are an abundant resource generated continuously by industry and human metabolism. Despite its high energy content, organic carbon is typically degraded to CO<sub>2</sub> through energy-intensive processes due to its heterogeneity, its low concentration, and stringent requirements for effluent quality. However, valorizing waste organics alongside recovered water is critical for the viability of utilities, industry, and next generation biorefineries. To that end, we employ a guantitative sustainable design (QSD) methodology to set a research agenda for the development of anaerobic membrane bioreactors (AnMBRs) for the conversion of dilute, aqueous organic carbon into methanerich biogas, with the simultaneous recovery of guality water. 150 unique AnMBR configurations were assembled as the landscape of possible development pathways. Each configuration was evaluated by integrating full-scale design and operation with techno-economic analysis (TEA) and life cycle assessment (LCA) in a Monte Carlo framework. Costs and environmental impacts were most sensitive to membrane configuration, membrane type, and inclusion of granular activated carbon (GAC) as physical media for membrane scouring. The least expensive designs (20th percentile) were exclusively AnMBRs with cross-flow, multi-tube membranes. Research targets were set through sensitivity analyses, prioritizing a decrease in cross-flow velocity (<0.5 m s<sup>-1</sup>), elimination of gas sparging, increase in membrane life (>10 years), decrease in upflow velocity for physical media bed expansion (<7.5 m  $h^{-1}$ ), and the development of low-cost physical media for fouling mitigation. Lastly, a subset of AnMBR designs had costs below state-of-the-art treatment (high rate activated sludge with anaerobic digestion). demonstrating the valorization of waste organics would be financially advantageous to industry and utilities.

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## Broader context

Aqueous waste organics are an underutilized energy resource due to their heterogeneous and dilute nature, but their valorization is essential to the financial viability and environmental sustainability of utilities and the biofuels industry. Here we demonstrate an emerging technology, the anaerobic membrane bioreactor (AnMBR), has the potential to reduce costs and key environmental impacts as compared to conventional aerobic technology. Through a quantitative sustainable design framework, we prioritize research objectives and set technology targets that will enable the financially viable production of methane and high quality water from dilute waste streams. Analyses of this nature are imperative to set agendas for research and technology development that have the greatest potential to valorize waste streams, and to align environmental and economic performance of technologies along more sustainable trajectories.

# 1 Introduction

Aqueous waste organics represent a ubiquitous, renewable resource for bioenergy production. To date, aerobic degradation of organics to carbon dioxide ( $CO_2$ ) has been the primary approach to wastewater management, despite incurred energy costs of 1000–2000 kW h kg carbon<sup>-1.1-5</sup> As a result, wastewater treatment plants (WWTPs, a.k.a. water resource recovery facilities,

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WRRFs) consume an estimated 3% of the US electricity demand, or 15 GW in the U.S. alone.<sup>2,6</sup> Additionally, as next generation biorefineries come online, wastewater treatment will be responsible for roughly 20% of capital costs and parasitic losses on the order of 11–26% of a refinery's operational electricity consumption.<sup>7–9</sup> As utilities and governments invest hundreds of billions of dollars in wastewater infrastructure over the coming decades,<sup>10</sup> the valorization of industrial- and municipal-derived aqueous waste organics can play a key role in aligning economic and environmental performance along sustainable trajectories.<sup>6,11–13</sup>

Mainline anaerobic wastewater treatment has been shown to have several benefits over traditional aerobic processes, including the potential to eliminate aeration (often half of a WWTP's electricity consumption<sup>14</sup>), reduce sludge wasting, and convert organic carbon to usable fuel (e.g., methane, hydrogen, electricity).<sup>6,15-18</sup> Mainline anaerobic treatment has been slow to take hold in technologically advanced communities due to several critical challenges, including large land requirements, unreliable performance at low temperatures, the potential to produce odors, and poor effluent quality.<sup>18</sup> Methane-producing technologies - which demonstrate the highest levels of energy production (as kJ g carbon removed $^{-1}$ )<sup>17</sup> – also suffer from high concentrations of soluble methane in reactor effluent.<sup>18</sup> However, technological advances continue to improve soluble methane recovery,19 and long-term performance of psychrophilic (<20 °C) anaerobic technology has been demonstrated.<sup>20,21</sup> A persistent barrier to anaerobic technologies, however, has been their failure to achieve adequate organic carbon removal to reliably meet discharge requirements.<sup>17</sup> With the introduction of membranes to mainline anaerobic treatment, anaerobic membrane bioreactors (AnMBRs) overcome this critical barrier and have been demonstrated to consistently produce high quality effluent in conjunction with high levels of methane-rich biogas production.<sup>17,22-26</sup>

With significant potential to align goals for bioenergy production and wastewater treatment, the research, development, and deployment (RD&D) of AnMBR is actively pursued. A key challenge in AnMBR RD&D, however, is the many disparate, competing designs and a lack of understanding of their relative sustainability and the factors governing environmental and economic performance.

The fundamental characteristic of AnMBR is the integration of an anaerobic unit process [e.g., completely stirred tank reactor (CSTR) or anaerobic filter (AF)] with membrane filtration.<sup>27</sup> The inclusion of membranes in the biological process allows hydraulic retention time (HRT) to be reliably decoupled from solids residence time (SRT); this often results in much smaller plant footprints (compared to conventional anaerobic treatment) and can lower capital costs.<sup>28</sup> However, the inclusion of membranes may result in trade-offs such as increased energy demand due to permeate pumping and, more significantly, membrane fouling control (e.g., gas sparging).<sup>29,30</sup> Ultimately, AnMBR is still an emerging technology, and many novel designs are being evaluated in parallel.<sup>24,25,31-41</sup> With a broad range of possible configurations (including reactor structure, fouling mitigation strategy, membrane type, etc.), it is expected that a subset of these alternatives are bound to be inherently better than others in terms of economic viability and environmental impacts of full-scale installations. To date, however, configurations have only been analyzed one-at-a-time, 30,42-44 limiting the field's understanding of which RD&D pathways have the greatest potential to align trajectories toward environmental and economic sustainability.

The objective of this work was to evaluate the environmental and economic implications of the broad landscape of AnMBR designs to prioritize RD&D needs. To this end, discrete design decisions were identified and assembled to generate 150 unique AnMBR designs, including various combinations of reactor design, membrane design, and methane management (Fig. 1, Table ESI-1<sup>†</sup>).



**Fig. 1** Discrete design decisions evaluated as potential development pathways for AnMBR. Initial choices limited downstream alternatives, as indicated by black lines and brackets. Dissolved methane recovery and utilization options were available to all designs. This approach resulted in a total of 150 discrete AnMBR designs that were evaluated as potential development pathways. A full list of evaluated designs can be found in Table ESI-1.†

Given that the focus of this study was on the identification of technology development pathways rather than the existing performance of individual configurations, it was assumed that each configuration could (at present or with incremental improvements over time) eventually achieve adequate effluent quality to meet permit requirements for biochemical oxygen demand (BOD) and suspended solids (e.g., 30 mg  $L^{-1}$  and 30 mg  $L^{-1}$ , respectively). This assumption is supported by a range of AnMBR designs that have been demonstrated to achieve  $\geq$  90% COD removal (Table ESI-6†). Full-scale designs for all configurations were developed and assessed using a quantitative sustainable design (QSD) methodology<sup>11</sup> integrating life cycle assessment (LCA) and techno-economic analysis (TEA) in a Monte Carlo framework in MATLAB. Sensitivity analysis was also conducted to determine which input parameters have the greatest influence on environmental and economic metrics. Finally, the relative importance of individual design decisions to economic and environmental sustainability were quantified and targets for research and development were prioritized.

## 2 Results and discussion

#### 2.1 Impact of design decisions

Membrane type [hollow fiber (HF), flat sheet (FS), multi-tube (MT)], membrane configuration (submerged, cross-flow), and GAC inclusion had the greatest relative median impacts on both cost (Fig. 2; 0.72, 0.19, and 0.39) and GWP (0.56, 0.58, and 1.28).



**Fig. 2** Relative impact of design decisions on global warming potential (GWP) and life cycle costs (as net present value, NPV). Values were calculated by dividing the difference in median values for a given metric (*i.e.*, cost or GWP) between two configurations that only varied by one decision by the minimum of the two values. A value of 0.1, therefore, represents a 10% difference in NPV or GWP stemming from that individual design decision. Intersections of similarly colored lines are the median of both data sets, circles indicate 25th and 75th percentiles, and the ends of each line are the 10th and 90th percentiles. For membrane material, the cost of all materials was assumed to be the same, so there is no observed variation in NPV due to membrane material choice.

Consistent trends were observed for reactor type, with AFs being less expensive than CSTRs (on average 6.2%), and for membrane types used in cross-flow configuration, with MT membranes consistently having the lowest GWP (median value 81% lower). Other choices, however, were more nuanced. For example, including GAC always reduced GWP (by 58% on average) and, when used in conjuction with HF membrane modules, lowered the total cost by 25%. When GAC was used in conjunction with FS modules, however, the additional membrane tank volume (FS required more tankage per m<sup>2</sup> of membrane surface area) increased the cost by roughly 40%. Although GAC eliminates the need for gas sparging, the material itself is expensive (Table ESI-2<sup>†</sup>). When a large amount of GAC is needed – as in the case of FS AnMBRs (which require a larger membrane tank than HF units handling the same flow) - the cost of GAC may outweigh any environmental benefits. This finding highlights the need for the development of membrane modules specifically designed to leverage physical media (e.g., GAC) for fouling mitigation, while also reducing membrane tank volume.

Methane management (*i.e.*, deciding whether or not to include a degassing membrane, DM) was also high in its impact on GWP because for any given design, if a DM was not included, anywhere between 30–50% of the produced methane was lost in the effluent<sup>26,35</sup> and was assumed to result in fugitive emissions to the atmosphere as a potent greenhouse gas (GHG). The degassing membrane is an early-stage technology with anticipated improvements in efficiency,<sup>45</sup> but a range of soluble methane management technologies may have potential for dramatic reductions in GHG reductions if energy positive operation or the beneficial use of soluble methane is achievable.

## 2.2 Impacts of individual designs

After examining the relative TEA and LCA impacts of the six discrete decisions under uncertainty, the cumulative impacts of each of the individual 150 AnMBR designs were compared to each other and to the performance of a state-of-the-art activated sludge system for organic carbon removal consisting of high rate activated sludge (HRAS; the first step in the AB process) coupled with anaerobic digestion (AD; see detailed explanation in ESI†). No single design universally minimized cost and environmental impacts across all categories. In order to comparatively examine the full landscape of designs, the median values of the net present value, net energy, and life cycle environmental impacts for each design were normalized between 0 and 1 according to eqn (1):

$$N_{ij} = \frac{M_{i,j} - M_{i,\min}}{M_{i,\max} - M_{i,\min}} \tag{1}$$

where *M* is the median value (from 3000 trials) of metric *i* for design *j*, *N* is the normalized value, and  $M_{i,\min}$  and  $M_{i,\max}$  represent the minimum and maximum median values (across all 150 designs) for metric *i*, respectively (Fig. 3; actual values for the minimum and maximum across all designs are reported in Table ESI-4†). Designs were also compared to a HRAS + AD treatment process using established kinetic and design parameters (details in Section S4 of the ESI†).<sup>4</sup>



**Fig. 3** Parallel coordinate plot comparing 150 designs based on costs (as NPV), net energy demand, and life cycle environmental impacts. For each metric, values are normalized between 0 and 1 by taking the difference between a given value and the minimum and dividing it by the range of the data set. For example, a total cost normalized value of 0.5 means the design had a cost that was half way between the design with the minimum cost and the design with the normalized value of 0.5 means the design had a cost that was half way between the design with the minimum cost and the design with the maximum cost. Designs with the lowest cost (*i.e.*, exclusively those with cross-flow, MT membranes) are blue and those that included submerged membranes and GAC are red. Designs below the black horizontal line in Net Energy Demand are energy positive.

In general, designs in the 90th percentile of costs had similar environmental impacts to those that were in the 10th percentile, indicating that certain design decisions decouple costs from energy consumption and environmental impacts. Specifically, the most expensive designs were principally submerged FS AnMBRs with GAC, with high costs stemming from larger membrane tanks (and thus, more GAC) as compared to HF systems (Fig. 4). The elimination of gas sparging through the use of an AFMBR (*i.e.*, the use of physical media for fouling mitigation) had dramatic benefits across environmental impact categories due to reduced electricity consumption, which led to strong environmental performance for all configurations



**Fig. 4** Comparison of cost [net present value, \$million] to net energy consumption [kW h m<sup>-3</sup>] for all 150 configurations. Diamonds and squares represent cross-flow and submerged configurations, respectively. See Fig. 3 for an explanation of the color scheme. Configurations below the bolded line are energy positive.

leveraging GAC. In particular, AFMBR systems demonstrated the greatest level of energy positive performance, with an estimated net energy consumption of -0.09 to -0.06 kW h m<sup>-3</sup>; significantly better than conventional activated sludge WWTPs which consume roughly 0.3 to 0.6 kW h m<sup>-3</sup> for similar levels of COD removal.<sup>1,3</sup> The HRAS + AD system was 13-34% more expensive than the top performing configurations (i.e. cross-flow with MT) - primarily due to aeration and sludge wastage requirements - but also operated near energy neutrality and had the lowest eutrophication impact. Eutrophication of all designs was driven by direct aquatic emissions (i.e., the effluent), and superior HRAS + AD performance stemmed from increased biomass yield and the assimilation of N and P into new biomass, thereby removing more nutrients from the wastewater during treatment. The relative difference between HRAS + AD and AnMBR eutrophication would be dramatically reduced for waste streams with more dilute nutrients or with N and P recovery downstream of AnMBR (discussed in more detail below). Designs above the 90th percentile for most environmental impacts (excluding ozone depletion and GWP) were cross-flow FS units because of the increased mass of membrane material needed and the added energy consumption from higher pumping rates to achieve adequate cross-flow velocity. The worst performing designs (above the 90th percentile) for ozone depletion and global warming were those that used membranes made of PTFE, which has a disproportionately high environmental footprint (normalized impact per kg of membrane material) as compared to other membrane materials that do not require fluorination.

Designs in the 20th percentile for cost had a cross-flow, MT membrane (blue lines in Fig. 3), which also achieved near energy neutral performance (-0.02 to 0.01 kW h m<sup>-3</sup>; Fig. 4). Although the AFMBR technology did achieve more energy

positive performance, the cost of GAC put AFMBR at ~ 10–31% higher life cycle cost than cross-flow systems with MT membranes. Focusing on the 10% of designs with the lowest cost (which varied by \$5 million in NPV) resulted in far less variability in LCA impacts than the full spectrum of designs, but even within these designs the median GWP impacts varied from 0.10 to 0.85 kg CO<sub>2</sub> eq. m<sup>-3</sup> treated. Thus, although cost minimization would steer design away from the worst (from a life cycle environmental perspective) designs, differences in environmental impacts may still be significant (varying up to an order of magnitude in the Monte Carlo simulations).

# 2.3 Sensitivity to scenario, process, and technology assumptions

The sensitivity of total net present value and GWP (both on a per m<sup>3</sup> treated basis) to each of the 22 input variables (Table 2) were determined using Spearman's rank correlation coefficients (Fig. 5), as described in the Methods section. Cost was most sensitive to interest, flux, and membrane lifetime, with median correlation coefficients of -0.50 (*i.e.*, a moderate negative correlation), -0.38, and -0.29, respectively. These values were significantly different from a "dummy" variable that had no effect on the model outputs (*p*-values < 0.001, Wilcoxon rank sum test). Membrane cost had a moderate positive correlation (0.16). Certain input parameters only impacted specific designs (*e.g.*, cross-flow velocity for designs with a cross-flow configuration); correlations for these parameters were only calculated for the relevant designs.

GWP was most sensitive to flux, which was shown to have a moderate negative correlation (-0.48) stemming from the impact on membrane tank volume and membrane materials (increasing flux

would decrease these sources of environmental impacts). This was statistically different from the "dummy" variable (p < 0.001). The influent soluble substrate concentration had an elevated 90th percentile (0.37). AF AnMBRs with a cross-flow MT membrane or AFs with a submerged membrane had higher sensitivity to this parameter. The percent of methane partitioned to the biogas (*i.e.*, the percent that is not dissolved in the effluent) had a high 90th percentile (-0.87) because of methane's potency as a GHG. Sensitivity to specific gas demand (SGD) was high (0.89), because increased SGD led to increased electricity consumption and, ultimately, higher GWP.

## 2.4 Research, development & deployment prioritization

Arguably the greatest strength of AnMBR is its ability to reliably, and under a range of temperatures and loading conditions,<sup>38,46</sup> achieve high levels of carbon removal (90–99%)<sup>17,26</sup> from wastewater. As further research is conducted on this emerging technology, treatment efficacy is expected to improve in conjunction with decreasing costs.<sup>44</sup> In order to expedite this process, the work presented here examined 150 possible AnMBR designs in order to prioritize research and development needs; a summary of these priorities can be found in Table 1.

AnMBRs may continue to be developed as both CSTRs and AFs, with submerged and cross-flow configurations. A critical challenge for cross-flow systems is the need to improve pumping efficiency relative to membrane scouring.<sup>47</sup> MT designs outperform existing FS designs in this respect, and membrane geometries targeting >0.02 m s<sup>-1</sup> cross-flow velocity per m<sup>3</sup> h<sup>-1</sup> feed flow should be prioritized to reduce pumping. Even with increased efficiency in membrane geometry, a fundamental tension may still exist between fouling mitigation (which is improved through



**Fig. 5** Sensitivity of (A) cost as NPV and (B) GWP to 22 input parameters. Values were determined using the results from Spearman's rank correlation calculations for all 150 configurations. Negative values indicate an inverse correlation between the input parameter and the output value. Tails are the 10th and 90th percentiles. The color of the boxes correspond to the colors in Fig. 1, where colors relate to the component of the AnMBR that is directly impacted. Inputs were also categorized based on whether they impact all scenarios or are specific to the technology/process.

#### Table 1 TEA and LCA impacts of key design decisions and proposed research and development priorities



higher shear) and the stability and performance of anaerobic bioprocesses (which may be negatively impacted by higher shear). High shear has been shown to disturb syntrophic interactions among anaerobic microorganisms (likely caused by a buildup of fermentation products),<sup>48</sup> indicating that the implications of shear on hydrolysis, microbial community structure, and overall reactor performance need to be examined further. As a result, cross-flow velocities below 0.5 m s<sup>-1</sup> should be targeted and fouling mitigation studies should also track bioprocess performance (CH<sub>4</sub> production, carbon removal, volatile fatty acid build-up, *etc.*) under continuous, long-term operation.

A critical challenge for submerged systems also stems from fouling mitigation, with this analysis demonstrating the clear need to eliminate gas sparging and replace it with less energy-intensive processes. Possible development pathways include the use of physical media for scouring of the membrane surface<sup>34,35,49</sup> or membrane vibration.<sup>18,50,51</sup> Including GAC in an AnMBR has many benefits (e.g., decreased membrane fouling,<sup>35,52,53</sup> higher fluxes,<sup>52</sup> and removal of trace contaminants),<sup>35</sup> but its use needs to be further evaluated through long-term experimental studies to better quantify trade-offs between an increased flux and a potential decrease in membrane life (increasing maintenance costs). To the degree possible, research efforts should focus on physical medium selection and membrane geometry to achieve membrane cleaning without damage, and simultaneously target cheaper, smaller, and lower density particles<sup>49</sup> to reduce upflow velocity requirements for bed expansion

(*e.g.*, reduce upflow velocity to  $<7.5 \text{ m h}^{-1}$  while keeping HRT below 1.6 h). Research on membrane materials can further support this goal by continuing to target better resistance to fouling while simultaneously increasing useful life.<sup>26,49</sup> Increasing membrane material efficiency (*i.e.*, the ratio of membrane surface area to material volume) to over 2000 m<sup>2</sup> m<sup>-3</sup> will also contribute to lower cost, more efficient systems with reduced environmental impacts. Finally, although researchers have achieved long-term AFMBR operation (485 days, 0.8 m<sup>3</sup> reactor volume) without chemical cleaning, cleaning may ultimately benefit system performance and a method should be developed that does not harm organisms attached to the fluidized GAC.<sup>53</sup>

The release of dissolved methane to the atmosphere via reactor effluent is a persistent problem for AnMBRs that has significant impacts on energy recovery and life cycle GWP.<sup>1,54,55</sup> Existing approaches include the use of sweep gas<sup>18,19</sup> or vacuum extraction with a degassing membrane;45 the latter was examined in this study as well as in Pretel et al.<sup>30</sup> If DMs are ultimately able to recover the same amount of energy (as methane) than that which is required to operate the system (as electricity), these systems would reduce the life cycle GWP of a plant by more than half but at a 5% increase in NPV. These environmental benefits may also be achieved through alternative uses of soluble methane, including the production of chemicals (e.g., methanol)<sup>56–60</sup> or the facilitation of denitrification.<sup>61</sup> Ultimately, the criticality of soluble methane management and the breadth of potential solutions underscores the need to develop integrated solutions for WWTPs interested in AnMBR. Beyond methane,

# 3 Conclusions

This novel analysis of AnMBR examined 150 potential development pathways, and identifies a range of research and development needs for this emerging technology. In addition to those specific needs identified above (and in Table 1), research efforts should continue to focus on core elements of AnMBR that reduce costs without sacrificing treatment efficacy (e.g., increasing membrane flux,<sup>54</sup> decreasing HRT<sup>26</sup>), maintaining the goal of achieving adequate effluent quality at costs below conventional aerobic systems. There were no designs that had both the lowest costs and lowest environmental impacts, though designs below the 10th percentile for costs did have comparatively low environmental impacts. Most design decisions resulted in trade-offs, the navigation of which is necessary if AnMBRs are to become broadly applied at full-scale WWTPs.<sup>30,44</sup> Relative to the stateof-the-art WWTPs relying on mainline aerobic treatment processes (HRAS + AD or the first stage of the AB process coupled with AD), two subsets of configurations - (i) cross-flow with multi-tube and (ii) submerged hollow fiber with GAC - were less expensive and able to achieve energy neutral or positive treatment. If future research targets the design, operational, and material improvements discussed here, net energy positive treatment and life cycle environmental benefits (e.g., net GHG removal from the atmosphere through energy offsets) can be achieved. In the U.S., hundreds of billions of dollars will be spent over the next few decades to meet more stringent discharge requirements and update aging infrastructure.<sup>10</sup> In parallel, existing and emerging biomass-based industries - including the biological production of fuels and chemicals - will generate dilute organic carbon waste streams that must be managed. Utilizing AnMBRs instead of energy-intensive aerobic processes will not only reduce costs and environmental impacts, but will ultimately align goals for local and global environmental sustainability with financial viability in the management of aqueous carbon resources.

## 4 Methods

## 4.1 Design roadmap

To evaluate the broad range of AnMBR designs that could be pursued in the coming years, the following discrete decisions were considered (Fig. 1): reactor type, membrane configuration, membrane type, membrane material, soluble methane management, and methane processing method. The landscape of possible combinations of these decisions (*e.g.*, a submerged membrane unit could only have a hollow fiber or flat sheet membrane module, not a multi-tube module) was assembled based on the literature and industry experience. Ultimately, the overall costs and environmental impacts of 150 different AnMBR designs were compared (Table ESI-1<sup>†</sup>). Details of individual decisions are described below.

4.1.1 Reactor type and configuration. The two reactor types considered in this study were CSTRs<sup>25,31,38,39,41</sup> and AF reactors.<sup>4,5,28,41,47</sup> For CSTRs, the volume of the reactor was dictated by HRT whereas AF sizing was determined by the organic loading rate (OLR; Table 2 lists values used for reactor design). The impact of including a down-flow aerobic sponge filter (AeF)<sup>63</sup> downstream of the AF was also evaluated, where AeF sizing was also dictated by OLR. Each type of reactor could use one of two membrane configurations: submerged<sup>24</sup> or crossflow.<sup>25,32</sup> Submerged systems were designed as side-stream AnMBRs (as described by Liao and colleagues<sup>28</sup>) with two reactors in series: a biological reactor followed by a membrane tank. Membrane tanks were as small as possible to accommodate the required number of membrane units, which was dictated by flux and design specifications from GE,<sup>64</sup> Kubota,<sup>65</sup> and Pentair.<sup>66</sup> If the design HRT (Table 2) was greater than the HRT of the membrane tank, a preceding CSTR was designed to accommodate the difference. Cross-flow systems were housed in a building adjacent to the biological reactors.

The implications of including physical media (assumed to be granular activated carbon, GAC, based on recent work on this topic<sup>34,35,40</sup>) in submerged configurations – designated the anaerobic fluidized MBR (AFMBR)<sup>35</sup> – was also examined. If GAC was included, it was placed only in the membrane tank and recirculation pumping was increased to achieve adequate upflow velocity for full bed expansion of the GAC; headloss due to bed expansion was assumed to be negligible.<sup>49</sup> No GAC replacement was assumed (consistent with Kim *et al.*<sup>35</sup>) and no aerobic polishing (consistent with Smith *et al.*<sup>44</sup>) postmembrane filtration was included.

**4.1.2 Membrane type and material.** The three membrane types considered in this study were hollow fiber (HF),<sup>24,31,35,37,40</sup> flat sheet (FS),<sup>25,36,38,41</sup> and multi-tube (MT).<sup>47,67</sup> While FS membranes can be used in either configuration, in full-scale systems HF or MT membranes are generally restricted to submerged or cross-flow configurations, respectively. In order to include specific membrane information (*e.g.*, nominal surface area per module) a default membrane was assumed for each type: GE ZeeWeed\* 500D for HFs, Kubota RM515 for FS, and Pentair X-flow for MT.

Three types of membrane material were included in the LCA: plastic, sintered steel, and ceramic. Within plastic, four types were considered: polyethersulfone (PES), polyvinylidenefluoride (PVDF), polyethylene terephthalate (PET), and polytetrafluoroethylene (PTFE).<sup>25,32,34,35,38,39,41</sup> Plastic was assumed to be usable for any membrane type, but sintered steel and ceramic were limited to FS and MT membranes, respectively. While these materials incur different environmental impacts due to their respective production methods, differences in costs were assumed to be negligible.

**4.1.3 Methane recovery and utilization.** At ambient temperatures, upwards of 50% of produced methane can be dissolved in AnMBR permeate (*i.e.*, reactor effluent).<sup>17</sup> If unutilized, dissolved methane would ultimately be lost to the atmosphere as a fugitive emission (*i.e.*, an unintended release to the natural environment), reducing bioenergy production and undermining broader

 Table 2
 Parameters varied during Monte Carlo simulation. For parameters with normal distributions, values shown are means with standard deviation in parentheses

Parameter	Units	Distribution	Values <sup>a</sup>	Citation
Flux	$L m^{-2} h^{-1}$	Triangular	5, 12, 17	36, 44 and 70
$TMP^{c}$	bar	Triangular	0.04, 0.17, 0.37	32, 33, 38, 44 and 70
Membrane life	years	Triangular	5, 10, 15	44
Membrane cost	$m^{-2}$	Uniform	64.5-107.6	71
Organic loading (AF)	$g \text{ COD } L^{-1} d^{-1}$	Uniform	0.2-8	4
Hydraulic loading (AF)	$m h^{-1}$	Uniform	2-6	5
Organic loading (AeF)	$g \text{ COD } L^{-1} d^{-1}$	Uniform	0.5-4	4 and 63
Hydraulic loading (AeF)	$m h^{-1}$	Uniform	0.11-0.44	72
HRT <sup>d</sup>	h	Uniform	8-12	36 and 44
HRT for $GAC^{e}$	h	Uniform	2.2-3.3	70
Recirculation ratio	Multiple of forward flow	Uniform	0.5-4	30 and 44
Cross-flow velocity	$m s^{-1}$	Uniform	0.4-2	28
Specific gas demand	$m^3 m^{-2} h^{-1}$	Uniform	0.05 - 1.2	30 and 44
Sparging frequency	%	Uniform	50-100	38
GAC concentration	$ m g~L^{-1}$	Uniform	187.8-225	35, 52 and 70
Biomass yield	g biomass g $COD^{-1}$	Uniform	0.02-0.08	16 and 44
Percent gaseous methane	%	Uniform	50-70	35 and 26
Upflow velocity for GAC bed expansion	$m h^{-1}$	Triangular	6, 8, 10	49
Flow rate	ML	Normal	75.7 (18.9)	Ь
Influent soluble substrate	$ m mg~COD~L^{-1}$	Normal	300 (30)	Ь
Influent particulate substrate	$mg$ COD $L^{-1}$	Normal	100 (10)	b
Interest	%	Uniform	6-10	b

<sup>*a*</sup> Probability density functions were characterized by: minimum and maximum values (uniform); minimum, most probable, and maximum values (triangular); average and standard deviation (normal). <sup>*b*</sup> Value was assumed. <sup>*c*</sup> Transmembrane pressure. <sup>*d*</sup> Hydraulic retention time. <sup>*e*</sup> Granular activated carbon.

goals to reduce global warming potential (GWP; methane is approximately 28 times worse than  $CO_2$  on a 100 year time horizon<sup>68</sup>). In order to recover all produced methane, the potential use of a degassing membrane (DM) was also included to quantify the trade-offs (in terms of cost and environmental impacts) of recovering methane or releasing it to the atmosphere.<sup>45</sup> Biogas produced and collected during the anaerobic process was assumed to be reused for energy and heat generation using a combined heat and power (CHP) system. Four CHP systems were evaluated in this study: internal combustion, combustion gas, microturbine, and fuel cell (their associated efficiencies can be found in Table ESI-2 in the ESI<sup>+</sup>).<sup>42,69</sup>

**4.1.4 Operation and cleaning.** Scouring of the membrane surface is necessary to remove foulants and varies depending on the reactor configuration, influent wastewater, and operational conditions.<sup>67</sup> Cross-flow systems utilized a high cross-flow velocity to mitigate fouling, whereas submerged AnMBRs relied on gas sparging. As an alternative to gas sparging, membrane scouring in submerged systems was also achieved by fluidizing a bed of GAC in direct contact with the membrane.<sup>35</sup>

For further foulant removal, chemical cleaning – both in-place (CIP) and out-of-place (COP) – may be required. Citric acid (100% by weight) and sodium hypochlorite (NaOCl, 12.5% by weight) were included in this study for inorganic and organic foulant removal, respectively. Annual consumptions were assumed to be 0.6 L year<sup>-1</sup> m<sup>-3</sup> d<sup>-1</sup> and 2.2 L year<sup>-1</sup> m<sup>-3</sup> d<sup>-1</sup> for citric acid and NaOCl, respectively, based on industry experience (Table ESI-2†). This assumption did not significantly impact the costs or life cycle environmental impacts of the system.

#### 4.2 Techno-economic analysis (TEA)

Using the system boundary described above, costs for the construction, operation, and maintenance of unit processes were calculated using equations derived from CapdetWorks<sup>TM</sup> (v3.0; Hydromantis Environmental Software Solutions, Inc.) and Hazen and Sawyer (Section S2 of the ESI†). In order to compare the costs of different configurations, the net present value (NPV) was calculated assuming an interest rate that varied uniformly between 6–10%. Unit costs for equipment were acquired from CapdetWorks<sup>TM</sup> 2014 vendor data or were retrieved directly from the manufacturer. See Section 2 of the ESI† for a detailed explanation of cost calculations.

### 4.3 Life cycle assessment

**4.3.1 Goal and scope definition.** LCA has been widely used to assess the broader environmental impacts of wastewater technologies and management decisions,<sup>73</sup> and was conducted here alongside net energy balance as an assessment of the environmental sustainability of design alternatives. LCA was carried out according to the methodology of ISO 14040/14044.<sup>74,75</sup> The functional unit for this study was the treatment of 1 m<sup>3</sup> wastewater (300 mg COD L<sup>-1</sup>) to discharge quality (30 mg COD L<sup>-1</sup>), with a project lifetime of 30 years. The system boundary included both construction and operation of the plant (Fig. ESI-1 in ESI<sup>†</sup>), but demolition was excluded (consistent with<sup>76–79</sup>). First and second order environmental impacts were also examined, where first order impacts were classified as direct emissions (to water, air, and land) from the WWTP and second order impacts stemmed from off-site processes such as

electricity production/transmission, material production, transportation, and avoided energy production offset by biogas recovery and utilization.

**4.3.2 Life cycle inventory (LCI).** The LCI for construction was generated from the detailed design of full-scale plants



**Fig. 6** Experimental data for key operational parameters for AnMBRs. Data were separated by membrane configuration (*i.e.*, submerged – square or cross-flow – diamond) and were labeled according to their citation (see Table ESI-6† for list of citations and values). If the configuration was listed as submerged, but a cross-flow velocity was given, the AnMBR was considered to have a cross-flow configuration. Data is presented as single points or a range of values, signified by error bars. Unfilled symbols represent pilot-scale AnMBRs (*i.e.*, >500 L); these were given preference when determining uncertainty distributions for the model (gray shading). Colors correspond to how the input affected the model (see Fig. 5 for a description). HRT has two distributions: one for AnMBRs with GAC (2.2–3.3 hours) and one for AnMBRs without GAC (8–16 hours).

using vendor specifications (*e.g.*, in the case of membrane modules, cassettes, and piping layouts), industry standards,<sup>4</sup> and multiplicative factors as described below. Once reactors, membrane cassettes, piping networks, and buildings (to house pumps, blowers, and cross-flow membrane units) were designed, the following items were quantified: volume of concrete, volume of excavation, mass of piping material, mass of membrane material, and mass of materials for the combined heat and power (CHP) system. A multiplicative approach was utilized to account for other miscellaneous WWTP construction materials (reinforcing steel, transportation, wiring, *etc.*) as outlined by Fahner<sup>80</sup> and Doka<sup>81</sup> and used by Foley and colleagues,<sup>82</sup> excluding those materials and processes that were directly quantified from detailed design (listed above).

For operation, considered processes included citric acid and NaOCl consumption for membrane cleaning, electricity consumption and offsets, GAC, membrane replacement, and sludge landfilling. Direct emissions from the plant to air and water were also included in this study (*i.e.*, COD, NH<sub>3</sub>, NH<sub>4</sub><sup>+</sup>, organic nitrogen, total phosphorus, and PO<sub>4</sub><sup>3-</sup> to water; CH<sub>4</sub> and CO<sub>2</sub> to air). The electricity mix was set to the 2014 U.S. average,<sup>83</sup> namely: 39% hard coal, 27% natural gas, 19% nuclear, 7% hydroelectric, 4.1% wind, 1.5% biomass, and 1% petroleum. For all materials and processes, LCI data were gathered using ecoinvent v3.0<sup>84</sup> and surrogates were used for items without available inventory data (Table ESI-3†).

**4.3.3** Life cycle impact assessment (LCIA). The LCIA was conducted using the tool for the reduction and assessment of chemical and other environmental impacts (TRACI v2.1), which classifies and characterizes the impact of each raw material and emission (quantified in the LCI) across nine categories: stratospheric ozone depletion, global warming potential (GWP), tropospheric smog formation, acidification, eutrophication, human health (HH) cancer, HH non-cancer, HH criteria-related (focusing on impacts of particulate matter and its precursors), and ecotoxicity.<sup>85</sup> No normalization, weighting, or aggregation beyond TRACI was performed.

#### 4.4 Uncertainty and sensitivity analyses

Monte Carlo with Latin Hypercube Sampling (LHS; 3000 trials per AnMBR design) was used to propagate input uncertainty for 22 parameters (Table 2) to quantify costs and life cycle environmental impacts for each design. The analysis was repeated with 1000 trials per design to confirm results did not change and that 3000 trials was more than sufficient to generate reproducible results. Assigned values were based on the literature or were conservatively estimated if data were lacking. Uncertainty distributions were also assigned to each parameter based on data availability; uniform distributions (*i.e.*,  $\pm 20\%$  of the assigned value) were used unless evidence suggested otherwise (Fig. 6). The sensitivity of net present value and GWP (both normalized to a per m<sup>3</sup> treated basis) to the input parameters across all configurations was determined by comparing Spearman's rank correlation coefficients.<sup>12,86</sup>

In order to discern how individual design decisions affected the model outputs, the relative impact of each decision was used to characterize its economic and environmental importance. Variation for a given metric (focusing on NPV and GWP) resulting from the six design decisions examined in this analysis was quantified by taking the difference in median values (*i.e.*, median of 3000 Monte Carlo trials) between two configurations that only varied by one decision (*e.g.*, the reactor type and membrane material were the same, but one used a submerged membrane and the other used a cross-flow), and dividing by the minimum of those two median values.

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# References

- 1 P. L. McCarty, J. Bae and J. Kim, *Environ. Sci. Technol.*, 2011, 45, 7100–7106.
- 2 USEPA, Wastewater management fact sheet: Energy conservation, EPA 832-F-06-024, 2006.
- 3 USEPA, Clean energy opportunities in water & wastewater treatment facilities: Background and resources, 2009.
- 4 G. Tchobanoglous, H. D. Stensel, R. Tsuchihashi and F. L. Burton, *Wastewater engineering: Treatment and resource recovery*, McGraw-Hill Higher Education, New York, London, 5th edn, 2013.
- 5 C. Grady Jr, G. T. Daigger, N. G. Love and C. D. M. Filipe, *Biological Wastewater Treatment*, CRC Press, 3rd edn, 2012.
- 6 B. E. Logan and K. Rabaey, Science, 2012, 337, 686-690.
- 7 R. Davis, L. Tao, E. Tan, M. Biddy, G. Beckham, C. Scarlata, J. Jacobson, K. Cafferty, J. Ross, J. Lukas, D. Knorr and P. Schoen, Process design and economics for the conversion of lignocellulosic biomass to hydrocarbons: Dilute-acid and enzymatic deconstruction of biomass to sugars and biological conversion of sugars to hydrocarbons, DOE Technical Report NREL/TP-5100-60223, 2013.
- 8 R. Davis, L. Tao, C. Scarlata, E. Tan, J. Ross, J. Lukas and D. Sexton, Process design and economics for the conversion of lignocellulosic biomass to hydrocarbons: Dilute-acid and enzymatic deconstruction of biomass to sugars and catalytic conversion of sugars to hydrocarbons, DOE Technical Report NREL/TP-5100-62498, 2015.
- 9 D. Humbird, R. Davis, L. Tao, C. Kinchin, D. Hsu, A. Aden, P. Schoen, J. Lukas, B. Olthof, M. Worley, D. Sexton and Dudgeon, Process design and economics for biochemical conversion of lignocellulosic biomass to ethanol: Diluteacid pretreatment and enzymatic hydrolysis of corn stover, DOE Technical Report NREL/TP-5100-47764, 2011.
- 10 ASCE, 2013 Report Card for America's Infrastructure, 2013.

- J. S. Guest, S. J. Skerlos, J. L. Barnard, M. B. Beck, G. T. Daigger, H. Hilger, S. J. Jackson, K. Karvazy, L. Kelly, L. Macpherson, J. R. Mihelcic, A. Pramanik, L. Raskin, M. C. M. Van Loosdrecht, D. Yeh and N. G. Love, *Environ. Sci. Technol.*, 2009, 43, 6126–6130.
- 12 X. Wang, P. L. McCarty, J. Liu, N.-Q. Ren, D.-J. Lee, H.-Q. Yu, Y. Qian and J. Qu, *Proc. Natl. Acad. Sci. U. S. A.*, 2015, **112**, 1630–1635.
- M. C. M. van Loosdrecht and D. Brdjanovic, *Science*, 2014, 344, 1452–1453.
- 14 WERF, Energy production and efficiency research The roadmap to net-zero energy, Water Environment Research Foundation Technical Report ENER1fs, 2011.
- 15 G. Tchobanoglous, F. L. Burton and H. D. Stensel, *Wastewater engineering: Treatment and reuse*, McGraw-Hill Professional, 2003.
- 16 B. E. Rittmann and P. L. McCarty, *Environmental biotechnology: Principles and applications*, McGraw-Hill, Boston, 2001.
- 17 B. D. Shoener, I. M. Bradley, R. D. Cusick and J. S. Guest, *Environ. Sci.: Processes Impacts*, 2014, **16**, 1204–1222.
- 18 D. J. Batstone and B. Virdis, *Curr. Opin. Biotechnol.*, 2014, 27, 142–149.
- 19 J. Cookney, E. Cartmell, B. Jefferson and E. McAdam, Water Sci. Technol., 2012, 65, 604–610.
- 20 C. Y. Gomec, J. Environ. Sci. Health, Part A: Toxic/Hazard. Subst. Environ. Eng., 2010, 45, 1169–1184.
- 21 A. L. Smith, S. J. Skerlos and L. Raskin, *Environ. Sci.: Water Res. Technol.*, 2015, 1, 56–64.
- 22 M. Dagnew, W. Parker, P. Seto, K. Waldner, Y. Hong, R. Bayly and J. Cumin, *Proceedings of the Water Environment Federation*, 2011, 2011, 4931–4941.
- 23 H. El-Mashad and R. Zhang, *Bioresour. Technol.*, 2010, **101**, 4021–4028.
- J. B. Giménez, A. Robles, L. Carretero, F. Durán, M. V. Ruano, M. N. Gatti, J. Ribes, J. Ferrer and A. Seco, *Bioresour. Technol.*, 2011, **102**, 8799–8806.
- 25 D. Martinez-Sosa, B. Helmreich, T. Netter, S. Paris, F. Bischof and H. Horn, *Bioresour. Technol.*, 2011, **102**, 10377–10385.
- 26 A. L. Smith, L. B. Stadler, N. G. Love, S. J. Skerlos and L. Raskin, *Bioresour. Technol.*, 2012, 122, 149–159.
- 27 C. Ramos, A. García and V. Diez, *Water Res.*, 2014, 67, 203–215.
- 28 B.-Q. Liao, J. T. Kraemer and D. M. Bagley, *Crit. Rev. Environ. Sci. Technol.*, 2006, 36, 489–530.
- 29 A. Hospido, I. Sanchez, G. Rodriguez-Garcia, A. Iglesias, D. Buntner, R. Reif, M. T. Moreira and G. Feijoo, *Desalination*, 2012, 285, 263–270.
- 30 R. Pretel, B. D. Shoener, J. Ferrer and J. S. Guest, Water Res., 2015, 87, 531–541.
- 31 L.-B. Chu, F.-L. Yang and X.-W. Zhang, *Process Biochem.*, 2005, **40**, 1063–1070.
- 32 J. Ho, S. Khanal and S. Sung, *Water Sci. Technol.*, 2007, 55, 79–86.
- 33 Z. Huang, S. L. Ong and H. Y. Ng, Water Res., 2011, 45, 705–713.
- 34 A. Hu and D. Stuckey, J. Environ. Eng., 2007, 133, 73-80.

- 35 J. Kim, K. Kim, H. Ye, E. Lee, C. Shin, P. L. McCarty and J. Bae, *Environ. Sci. Technol.*, 2011, 45, 576–581.
- 36 H. Lin, J. Chen, F. Wang, L. Ding and H. Hong, *Desalination*, 2011, **280**, 120–126.
- 37 A. Robles, M. V. Ruano, F. García-Usach and J. Ferrer, *Bioresour. Technol.*, 2012, **114**, 247–254.
- 38 A. L. Smith, S. J. Skerlos and L. Raskin, *Water Res.*, 2013, 47, 1655–1665.
- 39 I. Vyrides and D. Stuckey, Water Res., 2009, 43, 933-942.
- 40 R. Yoo, J. Kim, P. L. McCarty and J. Bae, *Bioresour. Technol.*, 2012, **120**, 133-139.
- 41 M. Harb, Y. Xiong, J. Guest, G. Amy and P.-Y. Hong, *Environ. Sci.: Water Res. Technol.*, 2015, 1, 800–813.
- 42 R. Pretel, A. Robles, M. V. Ruano, A. Seco and J. Ferrer, *Bioresour. Technol.*, 2013, **149**, 532–540.
- 43 R. Pretel, A. Robles, M. V. Ruano, A. Seco and J. Ferrer, *Sep. Purif. Technol.*, 2014, **126**, 30–38.
- 44 A. L. Smith, L. B. Stadler, L. Cao, N. G. Love, L. Raskin and S. J. Skerlos, *Environ. Sci. Technol.*, 2014, 48, 5972–5981.
- 45 W. M. Bandara, H. Satoh, M. Sasakawa, Y. Nakahara, M. Takahashi and S. Okabe, *Water Res.*, 2011, **45**, 3533–3540.
- 46 C.-H. Wei, M. Harb, G. Amy, P.-Y. Hong and T. Leiknes, *Bioresour. Technol.*, 2014, **166**, 326–334.
- 47 I. Martin-Garcia, V. Monsalvo, M. Pidou, P. Le-Clech, S. J. Judd, E. J. McAdam and B. Jefferson, *J. Membr. Sci.*, 2011, 382, 41–49.
- 48 S. I. Padmasiri, J. Zhang, M. Fitch, B. Norddahl, E. Morgenroth and L. Raskin, *Water Res.*, 2007, 41, 134–144.
- 49 M. Aslam, P. L. McCarty, J. Bae and J. Kim, Sep. Purif. Technol., 2014, 132, 10–15.
- 50 J. Kim, J. Shin, H. Kim, J.-Y. Lee, M.-h. Yoon, S. Won, B.-C. Lee and K. G. Song, *Bioresour. Technol.*, 2014, **172**, 321–327.
- 51 A. Kola, Y. Ye, P. Le-Clech and V. Chen, J. Membr. Sci., 2014, 455, 320–329.
- 52 R. Lee, P. L. McCarty, J. Bae and J. Kim, *J. Chem. Technol. Biotechnol.*, 2015, **90**, 391–397.
- 53 C. Shin, P. L. McCarty, J. Kim and J. Bae, *Bioresour. Technol.*, 2014, **159**, 95–103.
- 54 J. Gouveia, F. Plaza, G. Garralon, F. Fdz-Polanco and M. Peña, *Bioresour. Technol.*, 2015, **185**, 225–233.
- 55 S. Pacheco-Ruiz, S. Heaven and C. J. Banks, *Environ. Technol.*, 2015, **36**, 2328–2339.
- 56 E. Taher and K. Chandran, *Environ. Sci. Technol.*, 2013, 47, 3167–3173.
- 57 R. J. Conrado and R. Gonzalez, Science, 2014, 343, 621-623.
- 58 Q. Fei, M. T. Guarnieri, L. Tao, L. M. L. Laurens, N. Dowe and P. T. Pienkos, *Biotechnol. Adv.*, 2014, 32, 596–614.
- 59 C. A. Haynes and R. Gonzalez, *Nat. Chem. Biol.*, 2014, **10**, 331–339.
- 60 J. Sheehan, Nat. Biotechnol., 2009, 27, 1128-1129.
- 61 P. J. Strong, S. Xie and W. P. Clarke, *Environ. Sci. Technol.*, 2015, **49**, 4001–4018.
- 62 J. Delgado Vela, L. B. Stadler, K. J. Martin, L. Raskin, C. B. Bott and N. G. Love, *Environ. Sci. Technol. Lett.*, 2015, 2, 234–244.

- 63 M. Hatamoto, H. Yamamoto, T. Kindaichi, N. Ozaki and A. Ohashi, *Water Res.*, 2010, **44**, 1409–1418.
- 64 GE Power and Water, ZeeWeed 500D Module, 2013.
- 65 S.-H. Yoon, Membrane bioreactor processes: Principles and applications, CRC Press, 2015.
- 66 Pentair, X-Flow compact 33 ultrafiltration membrane, 2015.
- 67 S. Judd and C. Judd, *The MBR book*, Butterworth-Heinemann, Oxford, 2nd edn, 2011.
- 68 IPCC, Climate change 2013: The physical science basis. Contribution of Working Group I to the fifth assessment report of the Intergovernmental Panel on Climate Change, Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2013.
- 69 USEPA, RDC, ERG and CHPP, Opportunities for combined heat and power at wastewater treatment facilities: Market analysis and lessons from the field, 2011.
- 70 J. Bae, C. Shin, E. Lee, J. Kim and P. L. McCarty, *Bioresour. Technol.*, 2014, 165, 75–80.
- 71 L. Rupp, City of Ashland Oregon: Wastewater treatment plant membrane replacement options, Keller Associates Technical Report 210055/2/10-406, 2010.
- 72 A. Tawfik, R. A. Wahab, A. Al-Asmer and F. Matary, *Bioprocess Biosyst. Eng.*, 2011, **34**, 767–776.
- 73 L. Corominas, J. Foley, J. S. Guest, A. Hospido, H. F. Larsen,
   S. Morera and A. Shaw, *Water Res.*, 2013, 47, 5480–5492.
- 74 ISO, ISO 14040: Environmental management Life cycle assessment – Principles and framework, ISO Technical Report ISO 14040:2006(E), 2006.
- 75 ISO, ISO 14044: Environmental management Life cycle assessment – Requirements and guidelines, ISO Technical Report ISO 14044:2006(E), 2007.
- 76 J. M. Foley, R. A. Rozendal, C. K. Hertle, P. A. Lant and K. Rabaey, *Environ. Sci. Technol.*, 2010, 44, 3629–3637.
- 77 R. Nogueira, A. Brito, A. Machado, P. Janknecht, J. Salas, L. Vera and G. Martel, *Desalin. Water Treat.*, 2009, 4, 16–21.
- 78 N. Tangsubkul, P. Beavis, S. J. Moore, S. Lundie and T. D. Waite, *Water Resour. Manag.*, 2005, **19**, 521–537.
- 79 M. Yildirim and B. Topkaya, *Clean: Soil, Air, Water*, 2012, 40, 171–178.
- 80 S. Fahner, H. Bührer and C. Grabski, Thesis, IBB Ingenieurschule beider Basel, 1995.
- 81 G. Doka, Life cycle inventory of wastewater treatment, Swiss Centre for Life Cycle Inventories Technical Report Ecoinvent Report No., 13, 2003.
- 82 J. Foley, D. de Haas, K. Hartley and P. Lant, *Water Res.*, 2010, 44, 1654–1666.
- 83 U.S. Energy Information Administration, What is U.S. electricity generation by energy source?, 2014.
- 84 R. Frischknecht, N. Jungbluth, H.-J. Althaus, G. Doka, R. Dones, T. Heck, S. Hellweg, R. Hischier, T. Nemecek, G. Rebitzer and M. Spielmann, *Int. J. Life Cycle Assess.*, 2005, **10**, 3–9.
- 85 J. C. Bare, G. Norris, D. Pennington and T. McKone, J. Ind. Ecol., 2002, 6, 49–78.
- 86 S. Marino, I. B. Hogue, C. J. Ray and D. E. Kirschner, J. Theor. Biol., 2008, 254, 178–196.