1

2

3

# Cultivation of *Scenedesmus acuminatus* in different liquid digestates from anaerobic digestion of pulp and paper industry biosludge

Ran Tao, Aino-Maija Lakaniemi, Jukka A. Rintala

4 Laboratory of Chemistry and Bioengineering, Tampere University of Technology, P.O. Box 541, FI-

5 33101 Tampere, Finland

#### 6 Abstract:

7 Different undiluted liquid digestates from mesophilic and thermophilic anaerobic digesters treating 8 pulp and paper industry biosludge as such or after thermal pretreatment were characterized and 9 utilized for cultivating Scenedesmus acuminatus. Higher S. acuminatus biomass yields were 10 obtained in thermophilic digestates (without and with thermal pretreatment prior to anaerobic digestion (AD): 10.2 $\pm$ 2.2 and 10.8 $\pm$ 1.2 g L<sup>-1</sup>, respectively) than in pretreated mesophilic digestate 11 12  $(7.8\pm0.3 \text{ g L}^{-1})$ , likely due to different ammonium and sulfate concentrations in the digestates. S. 13 acuminatus removed over 97.4% of ammonium and 99.9% of both phosphate and sulfate from all 14 the digestates. Furthermore, color (74-80%) and CODs (29-39%) of the digestates were partially removed. The shown differences in methane yields (18–126 L CH<sub>4</sub> kg<sup>-1</sup> VS) from biosludge 15 16 resulting from the different AD processes and different microalgal yields emphasize the 17 importance of optimization of wood processing biorefineries and thus provide information to pulp 18 and paper industry development.

19

Keywords: wastewater treatment; pulp and paper industry; digestate characteristics; microalgal
growth; nutrient recovery

## 22 1 Introduction

23 Due to the environmental pollution and global warming, the European Council has promoted a 24 binding EU goal of greenhouse gas emissions with at least 40% internal reduction by 2030 compared to 1990, including 27% share of renewable energy for the EU (European Council, 2014). 25 26 Asia with the rapid growth and heavy dependence on fossil fuels (Lee et al., 2017) as well as other 27 regions e.g. North America, Latin America and Africa (Tan et al., 2017) must carry out a series of 28 policies and legislations for low-carbon and green growth. Biomass referring to all organic 29 materials that originate from plants (algae, trees and crops) can be converted into different kinds 30 of biofuels and energy carriers and is therefore one of the major renewable energy feedstocks (McKendry, 2002). Compared with other plants, microalgae have a high potential as a sustainable 31 32 bioenergy feedstock because of several advantages e.g. higher growth rate, no requirement for 33 arable land and potential for wastewater treatment to especially recover nutrients (Guldhe et al. 34 2017) but possibly also to remove toxic heavy metals (Romera et al. 2007) and remove faecal coliforms (Ansa et al. 2012). Besides, CO<sub>2</sub> from the exhaust gases of e.g. combustion, 35 metallurgical, chemical and biological processes can be utilized as carbon source for microalgal 36 37 cultivation (for a review, see Wang et al. 2008).

However, the existing problems, such as high demand for water and nutrients, low biomass production yields and high cost of microalgal harvesting, need to be solved before commercial utilization of microalgae to low-value products such as energy and fuels (Arenas et al. 2017). Since wastewater can provide the water and nutrients for the microalgae, many studies have been carried out to cultivate microalgae in different kinds of wastewaters including municipal, agricultural and industrial wastewaters (Ansa et al., 2012; Guldhe et al., 2017; Kinnunen and Rintala, 2016). 44 Microalgal cultivation in anaerobic digestion (AD) effluents, as a specific waste stream, has shown 45 significant potential for biorefinery applications due to the efficient nutrient removal and 46 accumulation of high-value products (e.g. astaxanthin, carotenoids and omega-3 fatty acids) to the 47 microalgal biomass (Polishchuk et al., 2015; Xia and Murphy, 2016). The integration of effluents 48 of AD from pulp and paper industry biosludge and microalgal cultivation (from now on referred 49 to as Integrated AD&MC system) has been studied to produce biomass and to recover nutrients 50 from wastewater (Kinnunen and Rintala, 2016; Polishchuk et al., 2015). The results of our previous study (Tao et al., 2017) indicated the possibility of high-yield microalgal biomass production and 51 52 efficient nutrient removal when *Scenedesmus acuminatus* was cultivated in liquid digestates from 53 AD of pulp and paper industry biosludge.

54 Pulp and paper industry is water and energy intensive biomass refining industry typically treating 55 its wastewaters in aerobic systems generating large amount of primary sludge and biosludge. 56 Anaerobic digestion of the generated sludges has gained increasing attention in pulp and paper 57 industry sludge treatment due to e.g. biomethane production as renewable energy (Kinnunen et al., 58 2015; Veluchamy and Kalamdhad, 2017) and possibility for nutrient recovery. More studies have 59 focused on anaerobic digestion of biosludge than primary sludge because primary sludge from 60 pulp and paper mill contains more wood fibres, which can be recycled to the fiber-processing 61 system of the mill instead of being anaerobically digested (de Alda, 2008; Kamali et al., 2016). 62 Biosludge also has a quite high content of lignocellulosic materials, which may limit its anaerobic 63 degradability (Kinnunen et al., 2015). To enhance biomethane production, application of pretreatment technologies have been considered. Thermal pretreatment prior to AD is one of the main 64 65 approaches used to enhance the methane production of pulp and paper industry biosludge (Kinnunen et al., 2015; Kamali et al., 2016). To understand the effect of thermal pretreatment 66

67 temperatures (80 °C, 105 °C, 121 °C and 134 °C) on methane production potential from biosludge 68 from pulp and paper industry Kinnunen et al. (2015) carried out biomethane potential batch assays 69 at 35 °C. They reported that biomethane production was increased by 39–140% compared to 70 untreated biosludge with the increasing pretreatment temperatures, except that biomethane 71 production from the biosludge treated at the lowest temperature, 80°C, was lower than that 72 obtained from the untreated one. However, although the increased pretreatment temperature 73 increased methane production, it also increases the costs and energy consumption of the thermal 74 pretreatment (Kinnunen et al., 2015). Because of this, Asunis (2015) further studied the anaerobic 75 digestion of pulp and paper mil biosludge at mesophilic and thermophilic conditions since the 76 operating temperature is a significant variable that also affects the methane yield. To our 77 knowledge, the AD plant reported to be under planning phase is the first full-scale AD plant 78 integrated in the pulp mill for digesting pulp mill sludges (Liikanen, 2016).

79 The previous studies show that biosludge with different treatments (pretreatment and AD 80 conditions) can result in different methane production yields and digestate compositions 81 (Kinnunen et al., 2015; Asunis, 2015). However, the microalgal cultivation in the effluents of AD 82 operated at different conditions has not been compared. Biomethane is generated during the AD 83 process while microalgal biomass can be produced during the cultivation by using the liquid 84 digestates from AD. To optimize Integrated AD&MC system for maximum bioenergy 85 (biomethane and microalgal biomass) production, it is important to study each process and thus 86 give an overview of the Integrated AD&MC system. The aim of this work was to study S. 87 acuminatus cultivation in various types of liquid digestates from AD of pulp and paper industry 88 biosludge and provide information with practical cases to biorefinery concept in pulp and paper 89 industries implementing AD and algal cultivation system simultaneously.

## 90 2 Materials and Methods

#### 91 2.1 Microalgal strain and liquid digestates

92 Scenedesmus acuminatus (SAG 38.81) was obtained from the SAG Culture Collection of Algae at 93 the University of Göttingen, Germany as a culture suspension. Stock culture was maintained in 94 100 mL N-8 medium in 250 mL Erlenmeyer flask on an orbital shaker (150 rpm) under fluorescent lamps (Osram L 18W/965 bio lux, Germany) at a light intensity of 40 µmol photos m<sup>-2</sup> s<sup>-1</sup>. The N-95 8 medium consisted of (g L<sup>-1</sup>): KNO<sub>3</sub>, 0.506; KH<sub>2</sub>PO<sub>4</sub>, 0.740; Na<sub>2</sub>HPO<sub>4</sub>, 0.260; MgSO<sub>4</sub>·7H<sub>2</sub>O, 96 97 0.050; CaCl<sub>2</sub>·2H<sub>2</sub>O, 0.018; FeNaEDTA·3H<sub>2</sub>O, 0.012 and micronutrient (ZnSO<sub>4</sub>·7H<sub>2</sub>O, 0.003; 98 MnCl<sub>2</sub>·4H<sub>2</sub>O, 0.013; CuSO<sub>4</sub>·5H<sub>2</sub>O, 0.018; Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>·18H<sub>2</sub>O, 0.007). An initial pH of 6.5 in the 99 N-8 medium was adjusted to 8.0 by adding 5 M NaOH.

100 Four types of digestates characterized in this study were collected from anaerobic semi-101 continuously fed completely stirred tank reactors (5 L liquid volume) treating biosludge from a 102 pulp and paper industry wastewater treatment plant (Asunis, 2015). The three different pulp and 103 paper mill biosludge digestates used in the microalgal cultivation experiments of the present study 104 were anaerobically digested at 55 °C (thermophilic digestate, T), anaerobically digested at 55 °C 105 after thermal pretreatment at 121 °C for 10 min (pre-treated thermophilic digestate, Tp) and 106 anaerobically digested at 35 °C after thermal pretreatment at 121 °C for 10 min (pre-treated 107 mesophilic digestate, Mp). The fourth pulp and paper mill biosludge digestate referred in this paper 108 was anaerobically digested at 35 °C (mesophilic digestate, M) (Asunis, 2015) and utilized for 109 cultivation of S. acuminatus in our previous study (Tao et al, 2017). The digestates were 110 centrifuged at 5200 rpm for 4 min, and the supernatant was filtered through a glass fibre filter 111 (Whatman GF/A, UK). After filtration, the liquid digestates were stored at 4 °C before use.

The microalgal growth results with the mesophilic digestate (M) are not directly comparable to the other digestates in the present study as *S. acuminatus* was grown in 1.5-times diluted mesophilic digestate M in the previous study (Tao et al., 2017), whereas in this study *S. acuminatus* was cultivated in undiluted digestates. Therefore, growth yields of *S. accuminatus* in digestate M are not compared to the microalgal cultivations results obtained in this study.

#### 117 2.2 Photobioreactors

118 S. acuminatus was grown separately in the three different digestates (digestate refers to liquid, 119 filtered digestate) for 21 days in photobioreactors (four replicates with each digestate), which 120 consisted of a 1-L glass bottle (PYREX) closed with a plastic cap with two tubes going through 121 the cap as the gas inlet and outlet. Air with 5% CO<sub>2</sub> (v/v) at a flow rate of 0.105 L min<sup>-1</sup> was 122 sparged from the bottom by a glass distribution tube (porosity 0, ø 22mm, Duran Group, Germany). 123 The photobioreactors were continuously illuminated using white fluorescent lamps (Osram L 18W/965 de lux cool daylight, Germany) with a light intensity of 240 µmol photos m<sup>-2</sup> s<sup>-1</sup>) from 124 125 two sides of the reactors. S. acuminatus was inoculated to the photobioreactors to provide an initial 126 optical density (OD) of 0.2. The initial total culture volume in the reactors was 600 mL. The 127 temperature of the reactors was maintained at 22±2 °C. Water evaporated during the cultivation 128 due to the constant sparging, and therefore distilled water was added to compensate the evaporated 129 water volume (marked with lines on the photobioreactors) each time before taking samples for 130 analyses.

#### 131 2.3 Analytical methods

The culture pH was measured using a WTW 330 pH meter (WTW, Germany) with a Slimtrode electrode (Hamilton, Germany). The light intensity was controlled by measuring the average value of six sites on two sides of the photobioreactors' outer surface by a MQ-200 Quantum Meter (Apogee, USA).

136 Volatile suspended solids (VSS) were measured by filtering 10–15 mL culture solution through a 137 glass fibre filter (Whatman GF/A). Each filter containing the suspended solids was dried at 105 °C 138 overnight, weighed and then burned in a 550 °C muffle furnace for 2 h and weighed again. VSS 139 was determined gravimetrically as a difference of the filters after treatment at these two 140 temperatures. The supernatant after VSS filtration was used in the analysis of digestate OD and 141 turbidity, soluble chemical oxygen demand (CODs), soluble biochemical oxygen demand (BOD<sub>7s</sub>), 142 dissolved organic carbon (DOC) and nutrients (N, P, S) concentrations. The OD was measured at 143 a wavelength of 680 nm using a Shimadzu UV-1700 Pharmaspec spectrophotometer after proper 144 dilution with distilled water to give absorbance values between 0.2–0.7. Turbidity was measured 145 with a TN-100/T-100 turbidimeter. OD was also measured from non-filtrated samples to assess 146 the microalgal biomass production.

147 CODs was determined using dichromate method according to the Finnish Standard SFS 5504. The 148 determination of BOD<sub>7</sub>s was done with a WTW OxiTop Control/ OxiTop measuring system. DOC 149 was measured with total organic carbon analyzer (Shimadzu Model TOC-5000) with ASI-5000 150 autosampler.  $NH_4^+$ -N was measured with an ion selective electrode (Thermo Scientific Orion ISE 151 meter). The potential extent of ammonium stripping was estimated by the following equation (Emerson et al. 1975) as rate of ammonia stripping has been shown to correlate well with unionized
ammonium concentration related to temperature and pH (Zimmo et al. 2003):

154 
$$unionized NH_3(\%) = \frac{100}{1 + 10^{(pK_a - pH)}}$$
 (1)

155 where  $pK_a = 0.09018 + \frac{2729.92}{T}$  and  $T = \text{temperature}({}^{\text{o}}\text{K})$ .

156  $NO_3^-$ ,  $NO_2^-$ ,  $PO_4^{3-}$  and  $SO_4^{2-}$  were measured using ICS-1600 ion chromatograph (Dionex, USA) 157 with AS-DV autosampler, Ion- Pac AS4A-SC anion exchange column, and ASRS-300 suppressor 158 (2 mm). The eluent contained 1.9 mM Na<sub>2</sub>CO<sub>3</sub> and 1.7 mM NaHCO<sub>3</sub>, and the eluent flow rate was 159 1 mL min<sup>-1</sup>.

## 160 **3 Results**

#### 161 3.1 Characteristics of the liquid digestates

162 The four pulp and paper industry biosludge digestates originating from digesters operated at 163 different temperatures treating biosludge with and without thermal pretreatment had different 164 characteristics (Table 1). The initial pH of all the digestates was above 8.0 and the high alkalinity 165 in the digestates provided a good buffering capacity for microalgal cultivation since S. acuminatus 166 prefers slightly alkaline conditions. The color of the digestates was measured by absorbance (OD) 167 at a wavelength of 680 nm and turbidity after removing the microalgal biomass by filtering. In 168 terms of OD, the color of the thermophilic digestates was higher than that of the mesophilic 169 digestates. In addition, OD value of the pretreated digestates was higher than those without 170 pretreatment. The digestate Tp showed the darkest color (OD: 0.63±0.08, turbidity: 320 NTU) of 171 all the digestates. However, the value of OD of the digestate T ( $0.59\pm0.06$ ) was higher than that of Mp (0.35±0.01) while the turbidity of the digestate T (280 NTU) was lower than that of Mp (290
NTU). Thus, there was no clear correlation between OD and turbidity.

The thermophilic digestates (T and Tp) had on an average 65 mg L<sup>-1</sup> higher ammonium 174 175 concentrations compared with the mesophilic digestates (M and Mp). In addition, the digestates treated at the same temperature resulted in 30–100 mg L<sup>-1</sup> higher ammonium concentration with 176 177 pretreatment than without pretreatment (Table 1). Ammonium was available in all the digestates 178 as nitrogen source for microalgal growth, while nitrate and nitrite concentrations were below 1.0 mg L<sup>-1</sup>. The total phosphorus content was similar (27-30 mg L<sup>-1</sup>) in all the digestates and 179 180 approximately 50% of the phosphorus existed in the form of phosphate except in the digestate M, 181 where phosphate share was slightly higher (64.3%). In addition, the sulfate-S concentration in 182 digestate Mp was much lower than that in the other three digestates (Table 1).

183 Similar phenomenon as with ammonium was observed with CODs values of the different 184 digestates. The thermophilic digestates had higher CODs values than the mesophilic digestates and 185 when the digestates produced at the same digestion temperature were compared, the biosludge 186 digestates generated in a process with pretreatment resulted in higher CODs than without 187 pretreatment (Table 1). The BOD<sub>7</sub>s/CODs ratios were lower than 1:20 in the measured digestates 188 (T, Tp and Mp), which means that most of the organic material left in the liquid digestates after 189 anaerobic digestion was not easily biodegradable. This indicates that the digestate could support 190 mainly photoautotrophic growth and that the microalgal growth in the digestates relied mainly on 191  $CO_2$  as the carbon source.

#### 192 3.2 Cultivation of *S. acuminatus* in the liquid digestates

#### 193 **3.2.1 Microalgal biomass production**

194 The microalgal biomass production as indicated by VSS in the three studied digestates (T, Tp and 195 Mp) was as shown in Fig. 1. The final microalgal biomass concentration after 21 days of batch cultivation was higher with both thermophilic digestates (T, Tp:  $10.2\pm2.2-10.8\pm1.2$  g L<sup>-1</sup>) than 196 that obtained with the mesophilic digestate (Mp:  $7.8\pm0.3$  g L<sup>-1</sup>). Despite the relatively high initial 197 ammonium concentrations (380-480 mg L<sup>-1</sup>) in all cultures, no clear lag phase was observed in 198 199 microalgal growth. The biomass concentration started to stabilize on day 15-18. In the beginning, 200 S. acuminatus in the digestate Tp grew slower than in the digestates T and Mp likely due to the 201 higher initial ammonium concentration and poorer light penetration (due to darker color of the digestate). Before day 9, S. acuminatus biomass concentration in the digestate T (6.0 g L<sup>-1</sup> VSS at 202 day 9) was the highest followed by *S. acuminatus* in the digestate Mp (4.9 g L<sup>-1</sup> VSS at day 9) and 203 Tp (4.4 g L<sup>-1</sup> VSS at day 9). Later, the culture concentration in the digestate Tp exceeded those in 204 205 the digestates Mp and T after day 9 and day 15, respectively.

#### 206

#### **3.2.2 Nutrient removal from liquid digestates**

S. *acuminatus* removed nutrients efficiently from the digestates (Fig. 2). The ammonium concentration decreased from initial 380–480 mg L<sup>-1</sup> to less than 0.2–10 mg L<sup>-1</sup>. The ammonium removal efficiency in the thermophilic digestates was over 99.9%, which was a little higher than the ammonium removal efficiency obtained in the mesophilic digestate (97.4%). The phosphate and sulfate were completely removed from all the liquid digestates in 7 days and 7–9 days, respectively. There was no clear difference in the phosphate and sulfate removal efficiency from the three different digestates (T, Tp and Mp).

#### 214 **3.2.3 CODs, DOC and color changes**

CODs removal efficiency was higher from the thermophilic digestates (38% and 39%) than from the mesophilic digestate (29%) (Fig. 3a). The DOC concentration in the thermophilic digestates decreased while DOC increased in the digestate Mp during the cultivation (Fig. 3b). The OD of the digestates was measured after removing the microalgae to show the digestate color change during the cultivation (Fig. 3c). The OD values in all digestates decreased until day 9 and remained quite stable after that. In the end of the batch cultivations, the color removal efficiencies in T, Tp and Mp were 80%, 74% and 79%, respectively.

#### 222 **4 Discussion**

223 This study shows that it is possible to produce a high concentration of microalgal biomass and 224 remove nutrients (ammonium, phosphate and sulfate) efficiently from various liquid digestates of 225 pulp and paper mill biosludge digestion without dilution. Thus, the results confirm the potential of 226 Integrated AD&MC system where the liquid digesates originate from pulp and paper wastewater 227 treatment plant biosludge digestion as shown in our previous study (Tao et al., 2017). It further 228 shows that the temperature of AD and thermal pretreatment prior to AD have some but not critical 229 impact on microalgal biomass production and nutrient, CODs and DOC removal. Thus, this study 230 provides information for decision making on the AD system to be invested simultaneously 231 considering the potential implementation of a microalgal system in a biorefinery concept in pulp 232 and paper industry. An overview (treatment methods of biosludge, microalgal cultivation 233 conditions and bioenergy production) of the studied Integrated AD&MC systems is shown in Table 234 2.

235 During the 21-day cultivation, approximately 35% more microalgal biomass (as VSS) was 236 obtained in the thermophilic digestates than in the mesophilic digestate. This is a promising 237 discovery, as methane production was also higher both without and with pretreatment in the 238 thermophilic digestion compared to the mesophilic process (Table 2) (Asunis, 2015) indicating 239 that the highest biogas production and microalgal biomass yield can be obtained in the same 240 integrated AD&MC system. However, it is clear that the influence from pulp and paper mill 241 digestate on microalgal growth is species specific as Kinnunen and Rintala (2016) previously 242 reported that biomass concentration less than 0.2 g  $L^{-1}$  (VSS) was obtained with a *Scenedesmus* 243 sp. originating from Lake Pyhäjärvi (Tampere, Finland) in an optimum mixture of 75% distilled 244 water and 25% liquid digestate from pulp and paper industry biosludge AD. Although the 245 biosludges used in Kinnunen and Rintala (2016) and this study were from the same pulp and paper 246 mill, the different characteristics of the digestates (likely due to the changes of e.g. wood source 247 used, pulp mill operation parameters and season) and microalgal strains clearly affect the 248 obtainable biomass quantity.

249 The effect of sludge pretreatment before digestion on microalgal cultivation is not, however, fully 250 clear based on this study. Asunis (2015) reported that thermal pretreatment increased the methane 251 yield by 100% in thermophilic AD process while the increase was 460% in mesophilic AD, while the methane yield with pretreatment at thermophilic condition was still 25 L CH<sub>4</sub> kg<sup>-1</sup> VS higher 252 253 than that obtained with pretreatment at mesophilic condition (Table 2). The difference caused by 254 the pretreatment prior to thermophilic digestion on the microalgal biomass production in the 255 digestate was not significant. Although highest methane and microalgal biomass production was 256 obtained at the same process (thermophilic AD with pretreatment), other factors should be 257 considered, including e.g. costs and energy burden of the thermal pretreatment and possible

removal of residual CODs from the digestate after microalgal cultivation. However, it is impossible to strictly compare the microlagal biomass yields in the mesophilic digestates with and without thermal pretreatment in this study due to the differences in dilution used in this study and our previous study (Tao et al, 2017).

262 The present study also shows that it is possible to obtain a high microalgal biomass yield in the 263 liquid digestates from pulp and paper wastewater treatment plant biosludge without dilution. This indicates e.g. that the digestate color levels in this study did not affect the microalgal growth greatly. 264 265 Substances in the liquid digestates causing the color may include clay, silt, finely divided inorganic 266 and organic matter, soluble colored organic compounds, and plankton and other microscopic 267 organisms (Wang et al., 2010). The turbidity of liquid digestates may vary ranging e.g. from 2960 268 to 51400 NTU as reported in the liquid fraction of mainly manure digestates from 11 full-scale co-269 digestion plants (Akhiar et al., 2017). The dark color of the medium is one of the issues, which 270 could reduce the microalgal growth due to poor light penetration (Wang et al., 2010; Marcilhac et 271 al., 2014; Xia and Murphy, 2016). For example, in a study by Wang et al. (2010), where Chlorella 272 sp. was cultivated in liquid fraction of anaerobically digested dairy manure (turbidity:1800–1900 273 NTU) with different dilutions (10, 15, 20 and 25-times) for 21 days the inverse correlation between turbidity and specific algal growth rates ( $R^2 = 0.982$ ) indicated that high turbidity can limit algal 274 275 growth. However, dilution for the microalgal growth increases total wastewater treatment volume 276 and might reduce microalgal growth owing to a reduction in nutrients and trace element 277 concentrations. In the present study, the different AD process conditions led to color differences 278 of the three liquid digestates (turbidity: 280–320 NTU), which however did not affect the color 279 removal efficiency. It is not easy to compare results (e.g. microalgal biomass production and 280 nutrients removal efficiencies) with other studies, which have similar medium turbidities as in the

present study since only few papers have mentioned the turbidity of medium during the microlalgalcultivation.

283 In this study, microalgal cultivation removed CODs to certain levels (29-39% removal) while DOC 284 acted somewhat contradictory to CODs as DOC level increased in the mesophilic digestate. COD 285 represents the demand of chemical oxidizer needed to oxidize all the oxidizable organic or 286 inorganic materials in wastewater and DOC is used to reflect dissolved organic carbon content of a sample. In most microalgal studies, either DOC or COD has been measured during microalgal 287 288 cultivation (Eloka-Eboka et al., 2017; Guldhe et al., 2017; Wang et al., 2010) and the correlation 289 between COD and DOC in microalgal cultures is not clear based on the previous studies. For 290 example, Marjakangas et al (2015) reported increase in both soluble COD and DOC concentrations 291 likely due to the low pH stress after C. vulgaris CY5 was mixotrophically cultivated in 292 anaerobically treated piggery wastewater. Thus, it seems that the changes in COD and DOC 293 depend on the growth conditions. In our study, organic carbon release from photosynthetic 294 microalgal cells might explain the observed increase in DOC during the cultivations in mesophilic 295 digestate. The decrease in CODs suggests that organic materials from the digestates were 296 consumed during the cultivation and the amount of consumed materials was higher than the 297 organic carbon released by the microalgae during normal photosynthetic growth. Some studies 298 have reported relatively high COD removal efficiencies (75-80%) from liquid digestates 299 integrated with microalgal cultivation (Yan and Zheng, 2014; Yang et al., 2015). The CODs in 300 digestes in this study was probably mainly caused by lignocellulosic materials (Kinnunen et al., 301 2015), which are not easily biodegradable and are therefore difficult to remove with biological 302 methods. Thus, further removal would be possible with other than biological treatments e.g. 303 chemical oxidation, if that would be deemed necessary.

304 The results in this study demonstrate that the microalgal cultivation efficiently removed 305 ammonium, phosphorus and sulfate from all digestates from pulp and paper industry biosludge 306 AD. Phosphorus was likely removed from the digestates through adsorption on the microalgal 307 surface, intracellular uptake and precipitation (Cai et al., 2013). In the present study, it is speculated 308 that enough phosphorus ensured microalgal growth since the microalgal biomass yield did not stop 309 or slow down when phosphate was no more detected from the liquid digestates after day 7. For 310 ammonium removal, several ammonium transformations (e.g. algal uptake, ammonia evaporation, 311 bacterial growth and nitrification) can occur in algae-bacteria consortium systems (González-312 Fernández et al., 2011). According to the average temperature (22 °C) and the observed pH range 313 (7.8-8.4), the theoretical fraction of unionized ammonia in all cultivations was 2.8%-10.3%. In 314 addition, low levels of nitrate were found in all cultivations. This data suggests that ammonium 315 stripping and nitrification may have occurred, but that the main portion of the removed ammonium 316 from the digestates was used for microbial growth.

317 Initial sulfate concentration in liquid digestates could affect the ammonium removal efficiency and 318 microalgal biomass production. This hypothesis is supported by the fact that the cultivations in the 319 digestates T and Tp having similar initial sulfate concentration (15-17 mg  $L^{-1}$ ) enabled over 99.9% 320 ammonium removal and similar microalgal biomass production while the different initial sulfate 321 concentrations in the digestates T and Mp (17 vs. 3 mg L<sup>-1</sup>), which had similar initial ammonium 322 concentration resulted in different ammonium removal efficiencies and algal biomass yield. 323 Biological nitrogen fixation is catalyzed during photosynthesis by nitrogenase, which contains 324 iron-sulfur clusters (Zheng and Dean, 1994) and the shortage of sulfur can therefore decrease the 325 assimilation of nitrogen (Kumaresan et al., 2017). Sulfate as primary sulfur source for microalgae 326 in aquatic environments has been proved to affect microalgal growth also in other studies (Ly et 327 al., 2017; Mera et al., 2016). Mera et al. (2016) reported that microalga *Chlamydomonas moewusii* 328 growth was quite similar at sodium sulfate concentrations 0.1-3 mM (SO<sub>4</sub><sup>2-</sup>-S:  $3.2-96 \text{ mg L}^{-1}$ ) but 329 microalgal biomass yields were lower at higher and lower sodium sulfate concentrations. In study 330 by Lv et al. (2017), similar *Chlorococcum* sp. growth at sulfate levels from 18-271 mg L<sup>-1</sup> was 331 obtained, but growth was much lower at 0 mg L<sup>-1</sup> sulfate.

## 332 **5 Conclusions**

333 Cultivatation of *Scenedesmus acuminatus* was succesful in different undiluted digestates from pulp 334 and paper industry biosludges treated at different AD conditions i.e. mesophilic vs. thermophilic, with and without thermal (121 °C) pretreatment. S. acuminatus grew well (7.8–10.8 g L<sup>-1</sup>) and 335 336 removed nutrients efficiently (over 97%) from all the digestates. Color (74-80%) and CODs (29-337 39%) were partially removed. The digestate from thermophilic process with pretreatment enabled highest microalgal biomass concentration, forming a promising discovery for pulp and paper 338 339 industry algae-based biorefinery applications as highest biomethane production was obtained at 340 the same conditions.

341

Acknowledgments: This work was supported by the Marie Skłodowska-Curie European Joint
Doctorate (EJD) in Advanced Biological Waste-To-Energy Technologies (ABWET) funded from
Horizon 2020 [grant number 643071]. We would like to thank Viljami Kinnunen and Ramasamy
Praveenkumar for the suggestions on the experimental set-up. We would also like to thank Tarja
Ylijoki-Kaiste for her help in the laboratory.

#### 347 **References**

348 1. Akhiar, A., Battimelli, A., Torrijos, M., Carrere, H., 2017. Comprehensive characterization of
349 the liquid fraction of digestates from full-scale anaerobic co-digestion. Waste Manage. 59, 118350 128.

- 2. Ansa, E.D.O., Lubberding, H.J., Gijzen, H.J., 2012. The effect of algal biomass on the removal
  of faecal coliform from domestic wastewater. Appl. Water Sci. 2, 87-94.
- 353 3. Arenas, E.G., Palacio, R., Juantorena, A.U., Fernando, S.E.L., Sebastian, P.J., 2017. Microalgae

as a potential source for biodiesel production: techniques, methods, and other challenges. Int. J.
Energy Res. 41, 761-789.

- 4. Asunis, F., 2015. Thermal pretreatment to enhance anaerobic digestion of pulp and paper mill
  biosludge (Unpublished master's thesis). Universita degli Studi di Cagliari Facolta di Ingegneria
  e Architettura, Italy and Tampere University of Technology, Finland.
- 5. Cai, T., Park, S.Y., Li, Y., 2013. Nutrient recovery from wastewater streams by microalgae:
  status and prospects. Renew. Sustainable Energy Rev. 19, 360-369.
- 361 6. de Alda, J.A.O., 2008. Feasibility of recycling pulp and paper mill sludge in the paper and board
  362 industries. Resour. Conserv. Recycl. 52, 965-972.
- 363 7. Eloka-Eboka, A.C., Inambao, F.L., 2017. Effects of CO<sub>2</sub> sequestration on lipid and biomass
- 364 productivity in microalgal biomass production. Appl. Energy 195, 1100-1111.
- 365 8. European Council. Conclusions (23 and 24 October 2014). 2030 Climate and Energy Policy
- 366 Framework. EUCO 169/14; 2014. Retrieved from
- 367 <u>http://www.consilium.europa.eu/uedocs/cms\_data/docs/pressdata/en/ec/145397.pdf.</u>

- 368 9. González-Fernández, C., Molinuevo-Salces, B., García-González, M.C., 2011. Nitrogen
  369 transformations under different conditions in open ponds by means of microalgae–bacteria
  370 consortium treating pig slurry. Bioresour. Technol. 102, 960-966.
- 371 10. Guldhe, A., Ansari, F.A., Singh, P., Bux, F., 2017. Heterotrophic cultivation of microalgae
  372 using aquaculture wastewater: A biorefinery concept for biomass production and nutrient
  373 remediation. Ecol. Eng. 99, 47-53.
- 374 11. McKendry, P., 2002. Energy production from biomass (part 1): overview of biomass.
  375 Bioresour. Technol. 83, 37-46.
- 12. Kamali, M., Gameiro, T., Costa, M.E.V., Capela, I., 2016. Anaerobic digestion of pulp and
  paper mill wastes–An overview of the developments and improvement opportunities. Chem. Eng.
  J. 298, 162-182.
- 379 13. Kinnunen, V., Rintala, J., 2016. The effect of low-temperature pretreatment on the
  380 solubilization and biomethane potential of microalgae biomass grown in synthetic and wastewater
  381 media. Bioresour. Technol. 221, 78-84.
- 14. Kinnunen, V., Ylä-Outinen, A., Rintala, J., 2015. Mesophilic anaerobic digestion of pulp and
  paper industry biosludge–long-term reactor performance and effects of thermal pretreatment.
  Water Res. 87, 105-111.
- 385 15. Kumaresan, V., Nizam, F., Ravichandran, G., Viswanathan, K., Palanisamy, R., Bhatt, P.,
- 386 Arasu, M.V., Al-Dhabi, N.A., Mala, K., Arockiaraj, J., 2017. Transcriptome changes of blue-green
- algae, *Arthrospira* sp. in response to sulfate stress. Algal Res. 23, 96-103.

- 16. Lee, C.T., Hashim, H., Ho, C.S., Van Fan, Y., Klemeš, J.J., 2017. Sustaining the low-carbon
  emission development in Asia and beyond: Sustainable energy, water, transportation and lowcarbon emission technology. J. Clean. Prod. 146, 1-13.
- 391 17. Liikanen, J. (2016, December 7). Biofuel from Äänekoski. Paper and Timber Journal.
- 392 Retrieved from <u>http://www.paperijapuu.fi/biofuel-from-aanekoski/</u>
- 18. Lv, J., Guo, J., Feng, J., Liu, Q., Xie, S., 2017. Effect of sulfate ions on growth and pollutants
  removal of self-flocculating microalga *Chlorococcum* sp. GD in synthetic municipal wastewater.
  Bioresour. Technol. 234, 289-296.
- 396 19. Marcilhac, C., Sialve, B., Pourcher, A.M., Ziebal, C., Bernet, N., Béline, F., 2014. Digestate
- color and light intensity affect nutrient removal and competition phenomena in a microalgal-bacterial ecosystem. Water Res. 64, 278-287.
- 399 20. Marjakangas, J.M., Chen, C.Y., Lakaniemi, A.M., Puhakka, J.A., Whang, L.M., Chang, J.S.,
- 400 2015. Simultaneous nutrient removal and lipid production with Chlorella vulgaris on sterilized
- 401 and non-sterilized anaerobically pretreated piggery wastewater. Biochem. Eng. J. 103, 177-184.
- 402 21. Mera, R., Torres, E., Abalde, J., 2016. Effects of sodium sulfate on the freshwater microalga
  403 *Chlamydomonas moewusii*: implications for the optimization of algal culture media. J. Phycol. 52,
  404 75-88.
- 405 22. Polishchuk, A., Valev, D., Tarvainen, M., Mishra, S., Kinnunen, V., Antal, T., Yang, B.,
- 406 Rintala, J., Tyystjärvi, E., 2015. Cultivation of Nannochloropsis for eicosapentaenoic acid
- 407 production in wastewaters of pulp and paper industry. Bioresour. Technol. 193, 469-476.
- 408 23. Romera, E., González, F., Ballester, A., Blázquez, M.L., Munoz, J.A., 2007. Comparative

- study of biosorption of heavy metals using different types of algae. Bioresour. Technol. 98, 3344-3353.
- 411 24. Tan, S., Yang, J., Yan, J., Lee, C., Hashim, H., Chen, B., 2017. A holistic low carbon city
  412 indicator framework for sustainable development. Appl. Energy 185, 1919-1930.
- 413 25. Tao, R., Kinnunen, V., Praveenkumar R., Lakaniemi, A.M., Rintala, A.J., 2017. Comparison
  414 of *Scenedesmus acuminatus* and *Chlorella vulgaris* cultivation in liquid digestates from anaerobic
  415 digestion of pulp and paper industry and municipal wastewater treatment sludge. J. Appl. Phycol.
  416 doi: 10.1007/s10811-017-1175-6
- 417 26. Veluchamy, C., Kalamdhad, A.S., 2017. Biochemical methane potential test for pulp and paper
  418 mill sludge with different food/microorganisms ratios and its kinetics. Int. Biodeterior.
  419 Biodegradation, 117, 197-204.
- 420 27. Wang, B., Li, Y., Wu, N., Lan, C.Q., 2008. CO<sub>2</sub> bio-mitigation using microalgae. Appl.
  421 Microbiol. Biotechnol. 79, 707-718.
- 422 28. Wang, L., Li, Y., Chen, P., Min, M., Chen, Y., Zhu, J., Ruan, R.R., 2010. Anaerobic digested
- 423 dairy manure as a nutrient supplement for cultivation of oil-rich green microalgae *Chlorella* sp.
- 424 Bioresour. Technol. 101, 2623-2628.
- 425 29. Xia, A., Murphy, J.D., 2016. Microalgal cultivation in treating liquid digestate from biogas
  426 systems. Trends Biotechnol. 34, 264-275.
- 427 30. Yan, C., Zheng, Z., 2014. Performance of mixed LED light wavelengths on biogas upgrade
- 428 and biogas fluid removal by microalga *Chlorella* sp. Appl. Energy 113, 1008-1014.

- 429 31. Yang, L., Tan, X., Li, D., Chu, H., Zhou, X., Zhang, Y., Yu, H., 2015. Nutrients removal and
- 430 lipids production by *Chlorella pyrenoidosa* cultivation using anaerobic digested starch wastewater
- 431 and alcohol wastewater. Bioresour. Technol. 181, 54-61.
- 432 32. Zheng, L., Dean, D.R., 1994. Catalytic formation of a nitrogenase iron-sulfur cluster. J. Biol.
- 433 Chem. 269, 18723-18726.

#### 434 **Figure Captions**

Fig. 1 Biomass concentration as volatile suspended solids (VSS) during the cultivation of *Scenedesmus acuminatus* in the liquid digestates from the pulp and paper wastewater treatment plant biosludge anaerobically treated at thermophilic process (55 °C) without pretreatment (T), with pretreatment at 121 °C for 10 min (Tp) and at mesophilic process (35 °C) with pretreatment at 121 °C for 10 min (Mp).

Fig. 2 The soluble ammonium-N (a), phosphate-P (b) and sulfate-S concentrations (c) during the cultivation of *Scenedesmus acuminatus* in the digestates from the pulp and paper wastewater treatment plant biosludges anaerobically treated at thermophilic condition (55 °C) without pretreatment (T), with pretreatment at 121 °C for 10 min (Tp) and at mesophilic condition (35 °C) with pretreatment at 121 °C for 10 min (Mp).

Fig. 3 CODs concentration and removal efficiency (a), DOC concentration (b) and OD of the cultivation medium (c), during the cultivation of *Scenedesmus acuminatus* in the digestates from the pulp and paper wastewater treatment plant biosludge anaerobically treated at thermophilic process (55 °C) without pretreatment (T), with pretreatment at 121 °C for 10 min (Tp) and at mesophilic process (35 °C) with pretreatment at 121 °C for 10 min (Mp).

450

451 **Tables** 

452 Table 1 Compositions of the liquid digestates from the anaerobic digestion of the pulp and paper industry
453 biosludge produced at thermophilic process without pretreatment (T) and with pretreatment at 121 °C for
454 10 min (Tp) and at mesophilic process without pretreatment (M) and with pretreatment at 121 °C for 10
455 min (Mp).

	Т	Тр	M <sup>a)</sup>	Мр
рН	8.2	8.3	8.5	8.3
Alkalinity <sup>a)</sup>	2700	3100	n.a.	2600
(mg L <sup>-1</sup> CaCO <sub>3</sub> )				
OD	$0.59\pm0.06$	$0.63\pm0.08$	$0.34\pm0.01$	$0.35\pm0.01$
Turbidity (NTU)	280	320	n.a.	290
$NH_4^+-N (mg L^{-1})$	$380 \pm 20$	$480 \pm 20$	$350 \pm 50$	$380 \pm 0$
$NO_{3}^{-}(mg L^{-1})$	<1.0	<1.0	<1.0	<1.0
$NO_2^{-}(mg L^{-1})$	<1.0	<1.0	<1.0	<1.0
TP (mg $L^{-1}$ )	$33 \pm 3$	$27 \pm 1$	$28\pm1$	$33 \pm 2$
$PO_4^{3-}-P (mg L^{-1})$	$16\pm3$	$15 \pm 3$	$18 \pm 1$	$15 \pm 1$
$SO_4^{2-}-S^{(a)} (mg L^{-1})$	$17 \pm 1.0$	$15 \pm 0.1$	$17\pm0.9$	$3 \pm 0.1$
CODs (mg L <sup>-1</sup> )	$1200\pm130$	$2000\pm130$	$910 \pm 30$	$1170\pm10$
$BOD_7s^{a)}(mg L^{-1})$	$110\pm5$	$60 \pm 100$	n.a.	$60\pm5$
DOC (mg L <sup>-1</sup> )	300±4	540±110	370±40	150±0

456 a) The values with  $\pm$  sign include standard errors

b) n.a.=data not available

Tuble I muchilite processes of underoole argestion of purp and puper made by prostadge and becheacesm
---

	Pretreatment	AD	Microalgal	Methane yield	Average microalgal
		temperature	cultivation	(L CH <sub>4</sub> kg <sup>-1</sup> VS)	biomass production
		(°C)	duration (d)		$(g L^{-1} VSS)$
М	No	35	14	18ª	8.8 <sup>b)</sup>
Мр	Yes	35	21	101 <sup>a</sup>	7.8
Т	No	55	21	63ª	10.2
Тр	Yes	55	21	126 <sup>a</sup>	10.8

459 *acuminatus* cultivation in the undiluted liquid digestates from the anaerobic digestion of the biosludge

460 a) data originated from Asunis (2015)

b) microalgae were cultivated in 1.5-times diluted digestate (Tao et al., 2017)

462

## **Figures**

**Fig. 1** 





**Fig. 2** 









