Development of a continuous flow reactor for aerobic granular sludge

By

Tanner Ryan Devlin

A thesis submitted to the Faculty of Graduate Studies of the University of Manitoba in partial fulfilment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

Department of Civil Engineering

University of Manitoba

Winnipeg

© Tanner Ryan Devlin 2019
To my family and friends.

Thank you
Merci beaucoup
Grazie mille
Wielkie dzięki
感激
خلي ممنون
Salamat
Acknowledgements

Dr. Oleszkiewicz for being the lead architect of my professional and personal development. You have carefully crafted and firmly cemented my ethos and values as a mentor, teacher, and friend.

Dr. Sparling for encouraging me to look beyond the stereomicroscope and down the *Francisella tularensis* hole. Also, for the raw materials to develop my applied microbiological skills. *Prost!*

Dr. Sears for providing exhaustive experience as a leading consultant in the field. As well, for reviewing documents more dry than an EAP report.

Doug and Lucy Devlin for patiently supporting me through my studies and life itself. None of this would have been possible without you and your unconditional love.

Charlene Urfano for always being supportive from one adventure to the next.

Micah Devlin for reminding me of my embarrassments and keeping me humble. Also, for the company during late-night study sessions.

*Towarzyszy broni,* Alessandro di Biase and Maciej Kowalski, for being the best of friends and colleagues. You were always able to deal with my *cazzate,* and you made our office a *casa.* I could not have asked for better *fratelli.*

Kevin Sagan and Marc de Rocquigny, thanks for being the best of friends and a welcome distraction from studies. You have been with me since the beginning.

Dr. Kruk and Dr. Jabari for paving the way before me and being role models. Dr. Wei and Dr. Yuan for always being open for discussion and warmly welcoming me to the University of Manitoba. Thanks to all who have been, are, and will be in the Environmental Laboratory.
Abstract

Aerobic granular sludge is an advanced biological process that provides sustainable wastewater treatment. Compared to conventional wastewater treatment processes, aerobic granular sludge exhibits better settling, higher volumetric loading, reduced chemical and electrical requirements, and less biosolids production. Currently, aerobic granular sludge is only available as a sequencing batch reactor process. Sequencing batch reactors, however, would not be suitable for retrofit at most existing wastewater treatment facilities. Therefore, many facilities would not be able to realize the benefits of aerobic granular sludge unless significant changes to existing infrastructure were made. A more compatible technology, being a continuous flow reactor, would facilitate the implementation of aerobic granular sludge into existing facilities that would otherwise struggle with sequencing batch reactors.

Thus, the goal of these studies was to develop a continuous flow reactor for aerobic granular sludge that would be suitable for retrofit into existing wastewater treatment facilities. To attain the goal, several items had to be addressed, including: 1) the suitability of aerobic granular sludge to treat low-strength, municipal wastewater; 2) the importance of hydrodynamic shear force for aerobic granular sludge stability; 3) the role of organic loading rate on aerobic granular sludge stability and performance; 4) the impact of selective wasting on granulation in continuous flow; and 5) the impact of reactor configuration on granulation in continuous flow. For more general items, such as suitability for low-strength wastewater and the role of hydrodynamic shear force, lab-scale sequencing batch reactors were used. The remaining tests were carried out in lab- and pilot-scale continuous flow reactors. Based on the results, design and operational parameters for an aerobic granular sludge-continuous flow reactor were developed and confirmed to be feasible for the treatment of low-strength, municipal wastewater.
# Table of contents

Acknowledgements ........................................................................................................................ iii

Abstract .......................................................................................................................................... iv

Table of contents ............................................................................................................................. v

List of equations ........................................................................................................................... xiii

List of figures ............................................................................................................................... xiv

List of tables ................................................................................................................................. xix

Nomenclature ............................................................................................................................... xxi

Executive summary ................................................................................................................... xxvii

Chapter 1: Introduction ................................................................................................................... 1

   Current and future trends in wastewater treatment ................................................................. 1
     Conventional wastewater treatment processes ................................................................. 1
     Commercially available options for upgrading existing treatment capacity ................. 2
     Case study: AquaNereda® sequencing batch reactor ................................................... 6
     Emerging energy efficient and sustainable nutrient removal technologies ............... 8

   Characteristics of AGS ........................................................................................................ 9
     Physical characteristics ...................................................................................................... 9
     Extracellular polymeric substances .............................................................................. 11
     Ion exchange and biologically induced precipitation ................................................... 13
Chapter 2: AGS-SBR treating low-strength wastewater at low hydrodynamic shear force........ 74

Introduction............................................................................................................................... 74

Methods..................................................................................................................................... 75

Reactor design and operation................................................................................................ 75

Analytical methods ............................................................................................................... 77

Results....................................................................................................................................... 80

Biomass characteristics and quantities.................................................................................. 80

Biomass growth characteristics............................................................................................. 84

Phosphorus removal .............................................................................................................. 86

Nitrogen removal .................................................................................................................. 87

Relationship between nutrient removal and biomass characteristics............................... 89

Summary ....................................................................................................................................... 92

Chapter 3: Impact of OLR on an AGS-SBR at low hydrodynamic shear force ...................... 96

Introduction............................................................................................................................... 96

Methods..................................................................................................................................... 98

Reactor design....................................................................................................................... 98
Chapter 4: Controlled SRT stabilizes AGS-SBR under high OLR and low hydrodynamic shear force

Introduction ............................................................................................................................. 117

Methods ................................................................................................................................... 119

Reactor design and operation .............................................................................................. 119

Analytical methods ............................................................................................................. 120

Results ..................................................................................................................................... 122

Reactor performance with SRT .......................................................................................... 122

Characteristics and morphology of biomass with SRT ...................................................... 126

Impact of stopping SRT ...................................................................................................... 129

Summary ............................................................................................................................. 133
Chapter 8: Impact of feast to famine ratios in an AGS-CFR

Introduction

Methods

Reactor configuration and operation

Source of inoculum

Wastewater characteristics

Analytical methods and calculations

Results

Reactor performance

Nitrogen and phosphorus profiles

Biomass characteristics and morphology

Summary

Bibliography

Chapter 9: Modelling the AGS-CFR with BioWin™

Introduction
Chapter 10: Engineering Significance .................................................................................. 210

Overview of conclusions ..................................................................................................... 210

Engineering significance .................................................................................................... 210

Low-strength wastewater .................................................................................................. 210

Hydrodynamic shear force ............................................................................................... 212

Solids retention time control at high organic loading rates .............................................. 213

Selective wasting in continuous flow reactors ................................................................. 214

Feast and famine conditions in continuous flow reactors ................................................. 216

Summary of findings ......................................................................................................... 217

Recommendations for future studies .................................................................................. 220

Appendix A ....................................................................................................................... 223

Determination of $D_F$ and $D_S$ ....................................................................................... 223

Appendix B ....................................................................................................................... 229

One-way ANOVA ................................................................................................................ 229

Appendix C ....................................................................................................................... 233

Determination of $r_{\text{max}}$ ............................................................................................... 233
Appendix D ........................................................................................................................................ 235

BioWin™ model of the AGS-CFR ............................................................................................... 235

Appendix E ...................................................................................................................................... 249

Hydrodynamic shear force versus airflow .................................................................................. 249
List of equations

Equation 1 ..................................................................................................................................... 79
Equation 2 ..................................................................................................................................... 79
Equation 3 ..................................................................................................................................... 79
Equation 4 ..................................................................................................................................... 79
Equation 5 ..................................................................................................................................... 79
Equation 6 ................................................................................................................................... 101
Equation 7 ................................................................................................................................... 101
Equation 8 ................................................................................................................................... 121
Equation 9 ................................................................................................................................... 121
Equation 10 ................................................................................................................................. 121
Equation 11 ................................................................................................................................. 121
Equation 12 ................................................................................................................................. 121
Equation 13 ................................................................................................................................. 121
Equation 14 ................................................................................................................................... 161
Equation 15 ................................................................................................................................... 183
Equation 16 ................................................................................................................................... 183
Equation 17 ................................................................................................................................... 183
Equation 18 ................................................................................................................................... 185
Equation 19 ................................................................................................................................... 185
Equation 20 ................................................................................................................................... 185
List of figures

Figure 1 Depiction of feast and famine phases by OUR profiles in a continuously aerated CAS-SBR (left) and continuously aerated AGS-SBR (right) with and without extended feast conditions due to high pCOD in the influent. The endogenous respiration phase in OUR profiles can be used to monitor the development and duration of the famine phase. ................................. 20

Figure 2 Estimated PAO to OHO ratio for a modelled biological treatment facility with increasing SRT and various configurations at a 12 h HRT. All models had typical primary influent characteristics of 500 mg-COD/L, 40 mg-TN/L, and 10 mg-TP/L as input. Fermentation was modelled at a 3 d SRT receiving primary sludge from a clarifier removing 70% of influent TSS............................................................................................................................................... 21

Figure 3 Steady-state MLSS and MLVSS in R1-R3. Error bars represent one standard deviation. ....................................................................................................................................................... 80

Figure 4 Steady-state TSS and VSS in the effluents of R1-R3. Error bars represent one standard deviation....................................................................................................................................... 81

Figure 5 Average diameter of granules from R1-R3. Error bars represent one standard deviation. ....................................................................................................................................................... 84

Figure 6 Left – steady-state whole sludge SRT in R1-R3; Right – steady-state Y_{net}. Errors bars represent one standard deviation....................................................................................................................................................... 85

Figure 7 Left – steady-state rate of OP release.; Right – steady-state rate of OP uptake. Errors bars represent one standard deviation. ....................................................................................................................................................... 87

Figure 8 Left – steady-state r_{max}, NOx-N prod. in R1-R3; Right – average TN removal, and the removal mechanisms, during steady-state. Error bars represent one standard deviation. .......... 88
Figure 9 Steady-state SND as a function of: MLVSS (top left); whole sludge SRT (top right); Y_{net} (bottom left); and R_{max, OP rel.} (bottom right). Errors bars represent one standard deviation. . 91

Figure 10 MLVSS for all AGS-SBRs over the entire study....................................................................................... 103

Figure 11 Specific COD, TN, and TP removal rates for each AGS-SBR during the last month of operation (i.e., steady-state). Error bars represent one standard deviation. ......................... 109

Figure 12 Percent sCOD utilized during the anaerobic influent distribution phase during a single reactor cycle. Negative values represent an increase in sCOD during the anaerobic feeding phase. ..................................................................................................................................................... 111

Figure 13 Mass of OP in solution for a single operational cycle on day 76. Anaerobic influent distribution occurred from 0-60 min and aeration was applied from 60-170 min. Initial values (i.e., t = 0 min) included the entire mass of OP supplied during the anaerobic influent distribution phase. Error bars represent one standard deviation. ................................................................. 112

Figure 14 Effluent COD, sCOD, TN, and TP in R1-R3. Error bars represent one standard deviation..................................................................................................................................... 122

Figure 15 sCOD profile during steady-state for R1-R3 over regular operational cycles. Error bars represent one standard deviation................................................................................................ 124

Figure 16 Nitrite/nitrate concentrations at the beginning and end of anaerobic influent distribution during steady-state for R1-R3. Error bars represent one standard deviation. .......... 126

Figure 17 Steady-state net observed yields shown with the nitrogen and phosphorus content of MLVSS. Error bars represent one standard deviation. ............................................................... 128

Figure 18 Stereoscopic images of biomass from R1-R3 during SRT control and seven days after SRT control was stopped. ........................................................................................................... 131
Figure 19 Top – Difference in COD, sCOD, TN, and TP removals in R1-R3 with and without SRT control (i.e., mixed liquor wasting); Bottom – Removal ratios (RR) for COD, sCOD, TN, and TP as a function of OLR. .............................................................. 132

Figure 20 Visual representation of filamentous microorganisms coating AGS under low hydrodynamic shear force. An increase in filamentous microorganisms coating AGS corresponded with the quantity of anaerobic COD uptake decreasing .......... 139

Figure 21 Cases of successful and unsuccessful granulation plotted with the OLR and USAV used in the respective study. ................................................................................. 140

Figure 22. AUSAV/A\textit{Nit} and AUSAV/A\textit{Het} as a function of SWL (top) and DO (bottom). ........... 141

Figure 23 Successful granulation with SRT control shown with the OLR limit without SRT control and the potential OLR limit with SRT control. It should be noted that OLR above 4.5 kg-COD/(m$^3$•d) were not examined, and therefore the OLR limit with SRT control may be higher than presented. ............................................................................................................................ 143

Figure 24 Visual representation of AGS stability with increasing OLR with and without SRT control at low hydrodynamic shear force ................................................................................. 143

Figure 25 Simplified schematic of the AGS-CFR in one of its embodiments ......................... 146

Figure 26 Steady-state MLSS and the ratio of MLVSS/MLSS with increasing selective pressure (i.e., rotational speed of the impeller in the clarifier). Error bars represent a single standard deviation ......................................................................................................................... 162

Figure 27 Average D$F$ and D$s$ of granules in the AGS-CFR with increasing selective pressure. Error bars represent a single standard deviation ......................................................................................................................... 164
Figure 28 Steady-state effluent characteristics for the AGS CFR under different selective wasting pressures. Higher selective wasting pressures were applied at higher rotational speeds of the impeller in the clarifier. Error bars represent one standard deviation.............................. 165

Figure 29 Steady-state concentrations of sCOD in the first and second aerobic compartment of the AGS-CFR under different rotational speeds for selective wasting. Error bars represent one standard deviation...................................................... 167

Figure 30 Steady-state anaerobic utilization of sCOD in the AGS-CFR under different rotational speeds for selective wasting. Error bars represent one standard deviation............................... 169

Figure 31 Ynet under different rotational speeds for selective wasting........................................ 171

Figure 32 Effluent TSS and VSS, and whole sludge SRT under different rotational speeds for selective wasting. Error bars represent one standard deviation................................. 172

Figure 33 Steady-state reactor nutrient profiles for configuration 1 and 2 without selective wasting. Error bars represent one standard deviation......................................................... 190

Figure 34 Top – steady-state SVI of configuration 1 and 2 without selective wasting (SW) and configuration 2 with selective wasting compared to inoculum. Error bars represented one standard deviation; Bottom – trend in SVI with the introduction of selective wasting to configuration 2................................................................................................................................. 196

Figure 35 Top – particle size analysis for configuration 1, configuration 2 without selective wasting, and configuration 2 with selective wasting; Bottom – total nitrogen removal for configuration 1 (i.e., steady state), configuration 2 without selective wasting (i.e., steady state), and configuration 2 with selective wasting (i.e., 12-48 d after selective wasting was started). Error bars represented one standard deviation......................................................................................................................... 199
Figure 36 Comparison of nutrient profiles between actual measured data and an optimized and unoptimized BioWin™ model.......................................................... 206

Figure 37 Relative abundance of select microbial guilds from the optimized BioWin™ model. .................................................................................................. 208

Figure 38 Visual representation of granulation as a function of selective pressure in CFRs. .... 214

Figure 39 Representation of existing activated sludge models (left) versus what components could be integrated with further studies into EPS production and modification in AGS (right). 221
List of tables

Table 1 Conventional technologies for increasing nutrient removal at existing CAS WWTPs [5]. ......................................................................................................................................................... 2

Table 2 Conventional technologies for increasing nutrient removal at existing CAS WWTPs [5]. ......................................................................................................................................................... 3

Table 3 Summary of expected MLSS concentrations with commercially available technologies for upgrading the capacity of CAS [9], [10], [12]–[14]........................................................................................................... 4

Table 4 Emerging process options for installing BNR within existing CAS footprints. ................. 9

Table 5 Summary of the main operating conditions affecting aerobic granulation..................... 30

Table 6 Summary of AGS-SBR pre-treatment configurations based on municipal wastewater characteristics....................................................................................................................................... 36

Table 7 Comparison of continuously-aerated AGS-SBRs to AGS-SBRs with anaerobic influent distribution. ....................................................................................................................................... 41

Table 8 Summary of design parameters for the AGS-CFR. ........................................................... 148

Table 9 Measured/estimated characteristics of equalized primary effluent from a full-scale domestic wastewater treatment plant. Values shown for steady-state operation of configuration 1 and configuration 2 with one standard deviation. Measured/estimated values were compared to default values in conventional modelling software [13].................................................................. 184

Table 10 Average steady-state effluent characteristics for configuration 1 and 2 shown with one standard deviation. .................................................................................................................. 187

Table 11 Summary of steady-state operational parameters for configuration 1 and 2 shown with one standard deviation. Target values indicated the planned values while estimated values indicated estimations from operational data. ...................................................................................... 194
Table 12 Comparison of select parameters between actual measured data and an unoptimized and optimized BioWin™ model.

Table 13 Summary of default and optimized BioWin™ parameters.

Table 14 Summary of operational parameters for AGS treating low-strength wastewater.
Nomenclature

AAO – anaerobic ammonium oxidizers

A/A – airflow to airflow ratio

AGS – aerobic granular sludge

A_{Het.} – airflow required to satisfy wastewater treatment without nitrification

A_{Nit.} – airflow required to satisfy wastewater treatment with nitrification

AOB – ammonium oxidizing bacteria

A_{USAV} – airflow required to satisfy the upflow superficial air velocity

bCOD – biodegradable chemical oxygen demand

BOD – biochemical oxygen demand

CAS – conventional activated sludge

CEPT – chemically enhanced primary treatment

CFR – continuous flow reactor

CMR – completely-mixed reactor

COD – chemical oxygen demand

DAMOA – denitrifying anaerobic methane oxidizing archaea

DAMOB – denitrifying anaerobic methane oxidizing bacteria

D_F – Feret’s diameter
DGAO – denitrifying glycogen accumulating organisms

DO – dissolved oxygen

DPAO – denitrifying phosphorus accumulating organisms

Dₚₚ – Diameter of equivalent sphere

EBPR – enhanced biological phosphorus removal

EPS – extracellular polymeric substances

F/M – food to microorganism ratio

ffCOD – flocculated and filtered COD

GAO – glycogen accumulating organisms

HRT – hydraulic retention time

HSR – high-strength reactor

IFAS – integrated fixed film activated sludge

LOT – limit of technology

LSR – low-strength reactor

MBfR – membrane biofilm reactor

MBR – membrane bioreactor

MEC – microbial electrolysis cell
MFC – microbial fuel cell

MLSS – mixed liquor suspended solids

MLVSS – mixed liquor volatile suspended solids

MSR – medium-strength reactor

N_{MLVSS} – nitrogen content of mixed liquor volatile suspended solids

NOB – nitrite oxidizing bacteria

OHO – ordinary heterotrophic organisms

OLR – organic loading rate

OP – orthophosphate

ORP – oxidation reduction potential

OUR – oxygen uptake rate

PAO – phosphorus accumulating organisms

pCOD – particulate chemical oxygen demand

PFR – plug-flow reactor

PHA – polyhydroxyalkanoate

P_{MLVSS} – phosphorus content of mixed liquor volatile suspended solids

PNA – partial nitritation with anammox

R – recycle ratio
RAS – return activated sludge

$r_{\text{COD}}$ – specific chemical oxygen demand removal

Re – Reynolds number

$r_{\text{max}, \text{NOx prod.}}$ – maximum specific nitrite/nitrate production rate

$R_{\text{max}, \text{OP rel.}}$ – maximum orthophosphate release rate

$r_{\text{max}, \text{OP rel.}}$ – maximum specific orthophosphate release rate

$r_{\text{max}, \text{OP up.}}$ – maximum specific orthophosphate uptake rate

RR – removal ratio

$r_{\text{TN}}$ – specific TN removal

$r_{\text{TP}}$ – specific TP removal

$r_{\text{WAS}}$ – specific wasting rate

S – concentration of substrate

SBR – sequencing batch reactor

$s_{\text{COD}}$ – soluble chemical oxygen demand

SND – simultaneous nitrification-denitrification

SOUR – specific oxygen uptake rate

SRT – solids retention time

SRTAGS – solids retention time of aerobic granular sludge
SRT_{sw} – solids retention time of selectively wasted biomass

SS – sidestream

sTN – soluble total nitrogen

sTP – soluble total phosphorus

SVI – sludge volume index

SWL – surface water level

TAN – total ammonia nitrogen

TIN – total inorganic nitrogen

TN – total nitrogen

TP – total phosphorus

TSS – total suspended solids

USAV – upflow superficial air velocity

v_{critical} – critical settling velocity

VER – volumetric exchange ratio

VFA – volatile fatty acid

VSS – volatile suspended solids

WAS – waste activated sludge

X – concentration of solids
$Y_{\text{net}}$ – net observed yield
Executive summary

The goal of these studies was to develop a continuous flow reactor for aerobic granular sludge that would be suitable for retrofit into existing wastewater treatment facilities. Several objectives were addressed to achieve the goal:

Suitability for low-strength wastewater. The suitability of AGS to treat low-strength wastewater was first examined on synthetic proteinaceous medium. With average influent of 340±30 mg/L COD, 42±5 mg/L TN, and 7±1 mg/L TP, an aerobic granular sludge-sequencing batch reactor achieved effluent orthophosphate of 0.2±0.4 mg-P/L and effluent total inorganic nitrogen of 17 mg-N/L. Aerobic granular sludge formed and matured, from conventional activated sludge inoculum, within 40 d while treating the synthetic proteinaceous medium. The aerobic granular sludge remained stable for the remaining 120 d experimentation period. Low-strength wastewater was used in all but one of the subsequent studies, and each of the studies successfully cultivated aerobic granular sludge whether it was on synthetic proteinaceous medium or primary effluent from a full-scale domestic wastewater treatment facility. The full-scale primary effluent averaged 310±60 mg/L COD, 48±9 mg/L TN, and 5.7±0.9 mg/L TP. Stable aerobic granular sludge did not loose performance despite chemical oxygen demand as low as 269 mg/L. Thus, granulation was both possible and stable while treating low-strength wastewaters such as primary effluent from domestic wastewater treatment facilities.

Importance of hydrodynamic shear force. The influence of increasing organic loading rate on aerobic granular sludge-sequencing batch reactor operation under low hydrodynamic shear force was examined. Specifically, upflow superficial air velocity was 0.41 cm/s while organic loading rate increased from 1.36 kg-COD/(m³•d) to 2.52 kg-COD/(m³•d) to 5.20 kg-COD/(m³•d). At 0.41 cm/s and 1.36 kg-COD/(m³•d), a mature aerobic granular sludge developed...
within 40 d. At organic loading rate of 2.52 kg-COD/(m³•d) a mature aerobic granular sludge developed within a month, but filamentous microorganisms were detected on granule surfaces approximately 15 d after mature-aerobic granular sludge development. Granules at organic loading rate of 2.52 kg-COD/(m³•d) were fully coated by a filamentous surface layer within two months from inoculation. At organic loading rate of 5.20 kg-COD/(m³•d) large granules developed rapidly, within two days, but then quickly destabilized. For the remaining four months of experimentation, biomass in the sequencing batch reactor with organic loading rate of 5.20 kg-COD/(m³•d) appeared as filaments joined together by a white, viscous substance that was likely a mixture of EPS and some cells. Thus, hydrodynamic shear force became more important as the organic loading rate increased, particularly when the organic load exceeded the capacity for anaerobic utilization. *It was therefore concluded that high hydrodynamic shear force was not a requirement for granulation, but higher shear may be beneficial at higher OLR.*

**Solids retention time as a control mechanism at high OLR.** An alternative method for stabilizing AGS, compared to high hydrodynamic shear force, was examined while treating brewery wastewater. Specifically, the controlled wasting of aerobic granular sludge, in addition to the selective wasting of slowly-settling biomass, was examined at organic loading rates from 1.5-4.5 kg-COD/(m³•d) and low upflow superficial air velocity of 0.51 cm/s. The measured whole sludge solids retention time was approximately 7 d under all organic loading rates. Previously, a minimum upflow superficial air velocity of 0.2 cm/s per kg-COD/(m³•d) was recommended to maintain stable aerobic granular sludge. Thus, at 4.5 kg-COD/(m³•d) the upflow superficial air velocity should have been 0.9 cm/s to generate stable aerobic granular sludge. However, it was demonstrated that controlled wasting of aerobic granular sludge helped maintain stable granules under lower than recommended upflow superficial air velocity. *Thus,*
controlling the solids retention time of aerobic granular sludge was effective at stabilizing the biomass at high organic loading rates and could potentially be used instead of hydrodynamic shear force.

**Impact of selective wasting.** The formation of aerobic granular sludge in a continuous flow reactor treating low-strength, proteinaceous wastewater (i.e., chemical oxygen demand = 370±30 mg/L; total nitrogen = 43±7 mg/L; and total phosphorus = 10±2 mg/L) under increasing selective pressure at organic loading rate of 1.5 kg-COD/(m³•d) was studied. A continuously-stirred clarifier was used to out-select the light fraction of biomass at 20 °C by operating at Re between 1500-5000. Successful granulation was observed, although below Re of 3000 a flocculent-granular biomass was developed and above Re of 4000 large granules with a filamentous population formed. Although mixed liquor suspended solids was low at Re number of 4000, granules had good characteristics (i.e., smooth surface, low filamentous microorganisms) and average diameter ranged from 0.21-0.67 mm. Average diameter at Re of 3000 only ranged from 0.21-0.23 mm. Thus, the optimal selective pressure was a Re between 3000-4000. *Therefore, selective wasting enhanced granulation in continuous flow, although too high a selective pressure could result in destabilization and undesirable biomass characteristics.*

**Reactor configuration.** The ratio of feast to famine conditions was examined in a continuous flow reactor consisting of multiple completely-mixed reactors in series. Two feast to famine ratios were examined: 1) 0.47 v/v; and 2) 0.1 v/v. Primary effluent from a full-scale municipal wastewater treatment plant was supplied to the aerobic granular sludge-continuous flow reactor and the organic loading rate was 0.7-0.8 g-COD/(m³•d). At lower feast to famine ratio, granulation was enhanced. Sludge volume index was 110±20 mL/g-VSS at feast to famine ratio of 0.47 while sludge volume was 86±8 mL/g-VSS at feast to famine ratio of 0.1. Granules
were also larger at lower feast to famine ratio. Average diameter ranged from 0.20-0.36 mm at 0.47 v/v, while average diameter ranged from 0.20-0.92 mm at 0.1 v/v. Selective wasting with critical settling velocity of 3.1 cm/min was integrated with the 0.1 v/v configuration. The SVI improved to 61±5 mL/g-MLSS and 50% of the granules were between 0.28-0.64 mm with selective wasting. *It was concluded that properly balanced feast and famine conditions enhanced the potential for granulation.*

Based on the results it could be concluded that aerobic granular sludge was suitable for continuous flow reactors. In fact, a full set of design parameters were developed and examined, demonstrating that an aerobic granular sludge-continuous flow reactor could treat low-strength, municipal wastewater to high effluent quality of <10 mg/L TIN and <1 mg/L TP.
Chapter 1: Introduction

Current and future trends in wastewater treatment

Conventional wastewater treatment processes

Water management is tied directly to the well-being of global populations and must be critically reviewed as municipalities improve infrastructure to meet future demand. By 2050 the world will contain nearly 10 billion people, of which a vast majority will reside in urban areas and experience an elevated standard of living [1], [2]. This will place severe strain on infrastructure and public resources, warranting the development and implementation of more sustainable practices to decrease the costs of incorporating advanced wastewater treatment processes. Currently, a majority of WWTP employ CAS to remove organic pollutants from wastewater with some nitrogen and phosphorus removed by biomass growth.

Slightly more of the influent TN and TP load can be removed by employing SS nitrogen and phosphorus removal technologies – Table 1. In most cases suitable effluent quality can be obtained by incorporating BNR into the existing CAS facility, simply by adding additional anoxic and anaerobic compartments. In environmentally sensitive areas, however, costly chemical addition and filtration technologies are required to meet the LOT [3], [4]. This results in high greenhouse gas emissions and large carbon footprints [5]. Thus, municipalities require processes that accomplish nutrient removal with minimal social and environmental impacts, as well as low financial burden — a “triple bottom line” approach [6]–[8].
### Table 1 Conventional technologies for increasing nutrient removal at existing CAS WWTPs [5].

<table>
<thead>
<tr>
<th>Technology</th>
<th>Process diagram of mainstream</th>
<th>Effluent quality</th>
<th>Relative Capex+Opex20 yr</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAS + SS-BNR</td>
<td><img src="image1" alt="Process diagram" /></td>
<td>TP &lt;3 mg/L&lt;br&gt;TN &lt;25 mg/L</td>
<td>1 (base case)</td>
</tr>
<tr>
<td>CAS-BNR LOT</td>
<td><img src="image2" alt="Process diagram" /></td>
<td>TP &lt;0.05 mg/L&lt;br&gt;TN &lt;2 mg/L</td>
<td>20</td>
</tr>
<tr>
<td>CAS-BNR</td>
<td><img src="image3" alt="Process diagram" /></td>
<td>TP &lt;1 mg/L&lt;br&gt;TN &lt;15 mg/L</td>
<td>10</td>
</tr>
</tbody>
</table>

**Commercially available options for upgrading existing treatment capacity**

Municipalities that are expanding often lack the footprint to install conventional nutrient removal technologies. Additional anoxic and anaerobic tanks, as well as the extended aeration capacity for ammonium oxidation, cannot be easily incorporated within existing footprints. Thus, new process options that retain more biomass in the existing system, thus achieving higher removal rates per unit volume, have gained popularity. There are several commercially available technologies that can increase the capacity of new and existing WWTPs – Table 2 and Table 3. Conventional systems that employ IFAS allow for increased capacity by decoupling SRT between the existing suspended growth culture and a newly integrated attached growth culture. Slower growing microorganisms, such as nitrifying AOB and NOB or AAO, become established in a dense biofilm on synthetic carrier material. This allows the suspended culture to be designed for faster growing heterotrophic processes such as carbon oxidation, facultative denitrification, and EBPR via PAO [9]. Without the addition of synthetic carrier material faster growing microorganisms would outcompete slower growing AOB and NOB, resulting in decreased treatment capacity of CAS at lower OLR than IFAS.
Table 2 Conventional technologies for increasing nutrient removal at existing CAS WWTPs [5].

<table>
<thead>
<tr>
<th>Technology</th>
<th>Process diagram</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFAS</td>
<td><img src="image" alt="IFAS Diagram" /></td>
<td>Synthetic carrier materials cultivate a dense biofilm of slow growing microorganisms, such as AOB and NOB, and are typically placed in the aerobic zone.</td>
</tr>
<tr>
<td>BioMag™</td>
<td><img src="image" alt="BioMag Diagram" /></td>
<td>Magnetite (i.e., SG = 5.2) combines with flocculent biomass, increasing the density of flocs and bolstering sedimentation of the entire system.</td>
</tr>
<tr>
<td>MBR</td>
<td><img src="image" alt="MBR Diagram" /></td>
<td>Membranes replace clarifiers, thereby ensuring effluent TSS of near 0 mg/L and a significant reduction in effluent particulate TN and TP.</td>
</tr>
<tr>
<td>Nereda®</td>
<td><img src="image" alt="Nereda Diagram" /></td>
<td>AGS is cultivated in a SBR with simultaneous influent distribution and effluent discharge, generating a dense biomass capable of EBPR and SND.</td>
</tr>
</tbody>
</table>

Synthetic carrier material is typically applied to the aerobic compartments of IFAS systems, although it has been applied in pre-anoxic compartments to increase capacity of denitrification. Carrier material has also been used in post-anoxic zones where the addition of external carbon source, such as methanol, was used for denitrification with low sludge yields [10]. The suspended growth culture may also be completely removed from the IFAS process by eliminating the RAS line and using more synthetic carriers, up to 67% of the reactor volume. Without RAS the IFAS system becomes a MBBR and the removal of contaminants is performed by the attached growth culture [11]. One disadvantage of commercially available MBBR systems is that phosphorus must be removed chemically since the carrier material cannot, yet, be cycled through an anaerobic phase. Since IFAS utilizes a “mobile” suspended culture that travels through different zones, a culture of PAO for EBPR can be selected for with the simple addition of an anaerobic compartment receiving raw wastewater, or primary fermentate rich in VFAs.
Table 3 Summary of expected MLSS concentrations with commercially available technologies for upgrading the capacity of CAS [9], [10], [12]–[14].

<table>
<thead>
<tr>
<th>Technology</th>
<th>MLSS</th>
<th>Carrier material</th>
<th>Solids separation</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAS</td>
<td>2-4 kg/m³</td>
<td>-</td>
<td>clarifier</td>
</tr>
<tr>
<td>IFAS</td>
<td>1-2 kg/m³ *</td>
<td>synthetic media</td>
<td>clarifier</td>
</tr>
<tr>
<td>BioMag™</td>
<td>5-10 kg/m³</td>
<td>magnetite</td>
<td>clarifier</td>
</tr>
<tr>
<td>MBR</td>
<td>4-10 kg/m³</td>
<td>-</td>
<td>membrane</td>
</tr>
<tr>
<td>Nereda®</td>
<td>4-15 kg/m³</td>
<td>-</td>
<td>settling period</td>
</tr>
</tbody>
</table>

*additional concentration of attached biomass integrated into existing CAS systems.

Ballasted sludge processes, such as BioMag™, are like IFAS and MBBR systems since they utilize a ballast to enhance capacity. Magnetite, a dense ferromagnetic material (i.e., SG = 5.2), is used in the BioMag™ process to promote a good settling biomass, consequently allowing for higher solids concentrations and smaller reactor sizes [12]. The overall increase in MLSS is slightly higher for BioMag™ than it is for IFAS due to the smaller volumetric requirements of magnetite compared to the low-density carrier material employed in IFAS. Furthermore, the hybrid magnetite-floc particles in BioMag™ move freely throughout the bioreactor and are not localized to specific reactor compartments like the carrier material applied in IFAS. Thus, the capacity of the entire microbial community, and not just specific microbial processes such as nitrification, are enhanced by BioMag™. However, the dense nature of magnetite also means that the mixing energy required for BioMag™ is higher than CAS, IFAS, or MBBR systems. The magnetite is also not fully recovered from WAS, thus replacement costs for virgin magnetite are incurred during operation.
Alternatively, MBRs use enhanced physical separation through membranes to allow for an increase in MLSS rather than utilizing carrier material for higher concentrations of attached biomass [13]. The excellent solids separation achieved by MBRs allow for increased capacity and improved effluent quality within existing footprints, as well as the potential for wastewater reuse applications [15]. Furthermore, MBRs can have lower capital costs than CAS systems when permits start to approach the LOT [16]. However, operational, maintenance, and replacement costs for MBRs are much higher than that of CAS systems, resulting in higher costs over time [17].

The Nereda® AGS process is a SBR based technology that has the potential to provide good quality effluent (i.e., TN < 5 mg/L; TP < 0.3 mg/L) within a significantly reduced footprint compared to CAS systems [18]. Most of the space reduction achieved by the Nereda® process is the result of energy efficient nutrient removal pathways occurring within a dense granular biomass [14]. Overall, with a higher capacity due to dense biomass and the lack of secondary clarifiers, AGS systems like the Nereda® process allow for up to 75% reduction in overall plant footprint [14]. Although IFAS, BioMag™, and MBRs provide potential to increase system capacity in confined spaces, their complexity brings along additional consequences. For example, compared to CAS systems IFAS, BioMag™, and MBR systems require more energy for aeration and BioMag™ can require up to three times more energy for mixing [19]. On the other hand, the Nereda® AGS technology has shown to achieve 30% savings in electricity for aeration alone when run in parallel with CAS systems [18]. The SBR operation of Nereda® simplifies the treatment process and does not require complex mechanical equipment for mixing, pumping, or solids separation.
Case study: AquaNereda® sequencing batch reactor

Research on AGS began at Delft University of Technology in 1993. The technology matured as an outcome of Dutch concerted research and development [20]. The final product was Royal HaskoningDHV’s Nereda® process, the first and only commercially available AGS technology. Installations of Nereda® have demonstrated significant improvements with regard to process stability, effluent quality (i.e., TN <5 mg/L; TP <0.3 mg/L), and energy savings (i.e., 30% to 60%) compared to traditional activated sludge processes [18]. Other advantages of the technology included lower capital and operational expenses, and a small physical footprint [21].

The first full-scale application of Nereda® began in 2005 with a retrofitted milk storage tank treating 250 m³/d of cheese factory waste in Ede, Netherlands [22]. The technology was scaled up for municipal wastewater treatment in 2008 with a 5,000 m³/d demonstration facility at the Gansbaai sewage treatment plant in South Africa [23]. Although only required to attain TN less than 15 mg/L and TP less than 10 mg/L, the Gansbaai installation reliably achieved effluent TN of 10 mg/L and TP of 3.2 mg/L [18]. Success at the Ede and Gansbaai facilities proved that AGS technology was feasible for full-scale applications of both industrial and municipal wastes. The Gansbaai demonstration in particular allowed applications of Nereda® to expand, starting with the design and construction of Epe WWTP, Netherlands in 2010-2011 [14].

The Epe WWTP has been operational since September 2011 and consists of inlet works with screens and grit removal, followed by three Nereda® reactors and effluent polishing via gravity sand filters [23]. At an average flow of 8,000 m³/d the facility has been able to achieve effluent TN of 4 mg/L and TP of 0.3 mg/L with only 60% of the original energy requirements (i.e., 3,500 kWh/d to 2,000-2,500 kWh/d after installation; Niermans et al., 2014).
Nereda® installations continue to push their own limits, and the largest operating design to date was commissioned in 2013 at the 100,000 m³/d Garmerwolde WWTP, Netherlands. The additional capacity gained from two 9,500 m³ Nereda® reactors is 30,000 m³/day, and the tank sizes are similar to the world’s largest SBR tanks. The Garmerwolde WWTP has passed the one-year performance test period to validate that combined effluent meets the target of less than 7 mg/L of TN and less than 1 mg/L of TP. Energy consumption of the Nereda® reactors has shown to be 50 to 60% more energy efficient than the existing AB-system that was commissioned in 2005 [18]. A retrofit currently underway in Ringsend WWTP, Dublin, Ireland will soon exceed the capacity of Garmerwolde and become the world’s largest Nereda® installation at 600,000 m³/d. This was no small feat as the Ringsend WWTP already boosts the world’s largest two-story SBR system.

The Nereda® technology has also shown potential for retrofitting existing continuous sludge systems after a successful demonstration at the 70,000 m³/d Frielas WWTP in Lisbon, Portugal [18]. The demonstration SBR was setup in one of the six existing CAS bioreactors and operated in parallel with the remaining CAS bioreactors [14]. Data from 2012 has indicated a 30% energy savings for aeration alone when compared with the existing system [24]. Combining this with the energy savings that granules bring by not using settling tanks, sludge recirculation pumps, and post-filtration units, the overall electricity demand was expected to be 50% of the existing CAS process [18]. However, the Nereda® technology cannot fully convert plug-flow based CFR into completely granular sludge-based processes.
Emerging energy efficient and sustainable nutrient removal technologies

Several promising technologies are under development for energy efficient and sustainable nutrient removal from wastewater. These include AGS-CFR, PNA, MBfR, and MEC based systems – Table 4. The development of an AGS-CFR would generate an option for municipalities with existing plug-flow, CAS systems to double or triple the hydraulic capacity of their existing process while also incorporating nutrient removal to <5 mg-TN/L and <0.3 mg-TP/L.

Anammox based systems, such as PNA, are an attractive option for mainstream nitrogen removal since they are near carbon independent and require significantly less energy for aeration [25], [26]. However, there are still significant issues with anammox cultures working effectively at temperatures below 10 °C, as well as competition for nitrite by NOB. A more mature technology that is just know gaining attention is MBfRs [27], [28]. Instead of relying on gas transfer kinetics through the mixed liquor, air or other gases are supplied directly through a membrane to an attached growth culture. In doing so, the energy losses associated with pumping air into solution to obtain a DO setpoint are removed. However, there remain problems with controlling biofilm growth, so much so that 80% of the energy saved from conventional aeration is put back into biofilm maintenance systems (e.g., scraping).

One of the least studied technologies for nutrient removal is the integration of MECs into CAS [29], [30]. By using MECs, electrically active biofilms cultivate the electrode material and exchange electrons directly with the electrical system. It is anticipated that by applying a small, alternating voltage gradient that the biofilm community could simultaneously oxidize ammonium and then remove nitrate hydrogenotrophically. However, this technology is the least studied out of the four presented and the concept has not yet been demonstrated.
Table 4 Emerging process options for installing BNR within existing CAS footprints.

<table>
<thead>
<tr>
<th>Technology</th>
<th>Process diagram</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF AGS</td>
<td><img src="image1" alt="PF AGS Diagram" /></td>
<td>AGS is cultivated in PL conditions by selective wasting and metabolic management or maximizing anaerobic carbon utilization. Organics, TN, and TP are removed without internal recycle lines required for CAS-BNR. An anammox based system removes TN from wastewater with low carbon to nitrogen ratios. When combined with CEPT, readily bioavailable substrate could be used for denitrification of residual nitrate in a pre-anoxic zone.</td>
</tr>
<tr>
<td>Anammox</td>
<td><img src="image2" alt="Anammox Diagram" /></td>
<td>An MBfR utilizes membrane diffusion to supply air, pure oxygen, or hydrogen gas directly to a biofilm. If air/oxygen and hydrogen gas were supplied to the membranes TN removal would occur in the biofilm while a PAO culture could be developed in suspension. Like the MBfR, but in the complete absence of oxygen, a MEC could be used for SND with alternating current. At the anode ammonium would be oxidized while at the cathode hydrogen gas generated by electrolysis would be used for denitrification. PAO could be in suspension.</td>
</tr>
<tr>
<td>MBfR</td>
<td><img src="image3" alt="MBfR Diagram" /></td>
<td></td>
</tr>
<tr>
<td>MEC</td>
<td><img src="image4" alt="MEC Diagram" /></td>
<td></td>
</tr>
</tbody>
</table>

Characteristics of AGS

Physical characteristics

The compact, tightly aggregated characteristics of AGS are responsible for most of the benefits AGS processes exhibit over conventional systems. For instance, an AGS-SBR can obtain comparable treatment to conventional CAS within 70% of the footprint [31]. Furthermore, a 60% reduction in energy requirements has been demonstrated at a full-scale AGS plant in the Netherlands. Average diameter of AGS has been reported between 0.2 and 5 mm, although granules as large as 23 mm have been observed in lab-scale SBRs [32]–[36]. The large size and high density of AGS allows for excellent settling properties and low SVI from 20-80 mL/g [37]. Values for specific gravity from 1.004 to 1.1 have been reported for AGS, further demonstrating that granules are denser than flocs, which typically have specific gravities from 1.002 to 1.006 [37]–[39].
Like any physical system, in situ settling velocities are affected by temperature, viscosity, and geometric properties. Large granules (i.e., average diameter = 2.3 ± 0.5 mm) from a lab-scale reactor had settling velocities of 84 m/h at 5 °C and 145 m/h at 40 °C while small granules (i.e., average diameter = 1.5 ± 0.3 mm) had settling velocities of 35 m/h at 5 °C and 63 m/h at 40 °C [40]. Specific gravity and settling velocities of aerobic granular sludge are both higher than conventional activated sludge systems. Granules settle rapidly and as discrete particles, resulting in a complete overhaul of clarification principles applied to flocculent sludge used in conventional treatment. Analysis of full-scale granular biomass indicated that PAO had a higher density than GAO leading to significantly higher settling velocities for PAO dominated granules [41].

Granules can be highly porous, allowing for sufficient mass transfer through the microorganism-EPS complex. Because of porosity, concentration gradients form and stratification of aerobic, anoxic, and anaerobic layers develop throughout granule depth and cycle time. This in turn leads to microbial diversity across granule depth, and thus stratification of nutrient removal pathways [42]. Mass transfer in AGS can be disrupted by an overgrowth of filamentous bacteria on the outer surface layer of granules when readily available substrate is present under aerobic conditions [43]. Particulate matter can also contribute to granule surface “fouling” by acting as a constant source of biodegradable substrate to microorganisms in the surface layer under aerobic conditions [44]. Furthermore, overproduction of EPS in highly loaded granular sludge reactors can result in reduced pore space and reduced mass transfer leading to continuous breakage of granules [45]. Any detriment to AGS porosity will directly affect overall performance since mass transfer to localized areas containing specialized microorganisms would be compromised.
**Extracellular polymeric substances**

EPS has been recognized as a key element that shapes and provides structural support for biofilms [46]. It has also been suggested that the EPS can act as a buffer layer for cells against harsh external environments, and that components of EPS are used as carbon and energy sources during starvation periods [47]. Gel-forming exopolysaccharides, in particular, were observed as major structural materials in AGS [48]. Specifically, granulan and alginate-like exopolysaccharide, both producing gel-like structures, have been proposed as the gel forming constituents of the granules.

The distribution of EPS (i.e., as proteins, polysaccharides, and lipids) and cells (i.e., both live and dead) has been studied in AGS [49]. The authors found that in acetate-fed granules protein and β-D-glucopyranose polysaccharides formed the core whereas the cells and α-D-glucopyranose polysaccharides accumulated in the granule outer layers. The role of individual EPS components for structural stability of phenol-fed aerobic granules has also been examined [50]. Selective enzymatic hydrolysis of proteins, lipids, and α-polysaccharides had a minimal effect upon the three-dimensional structural integrity of the granules. Conversely, selective hydrolysis of β-polysaccharides fragmented the granules. The β-polysaccharides were expected to form the backbone of a network-like outer layer with embedded proteins, lipids, α-polysaccharides, and cells to support the mechanical stability of granules.
Alginate-like exopolysaccharides in AGS were observed to have significantly higher poly-guluronic acid blocks than mannuronic acid blocks, while alginate-like exopolysaccharides from flocculent sludge had equal amounts of both chemical blocks [51]. These differences encouraged a continuous network structure via crosslinks of guluronic acid blocks in AGS. Consequently, the guluronic acid blocks were demonstrated to promote higher resistance of aerobic granules, compared to flocculent biomass, to breakage when subjected to deformation.

The broad role of proteins and polysaccharides in maintaining the structural stability of the aerobic granules has been investigated [52]. Hydrolysis of extracellular proteins in the EPS matrix of aerobic granules led to collapse of the EPS matrix, and subsequent disintegration of aerobic granules. These observations suggested that extracellular proteins would be essential for maintaining structural stability of AGS, more so than polysaccharides. A reduction in the protein to polysaccharide ratio in the EPS matrix has commonly been considered a sign of weakening AGS structure [53], [54]. Indeed, because of their simple molecular structure, polysaccharides are more sensitive to biological and chemical degradation than proteins. Therefore, granules with low protein to polysaccharide ratios have a matrix consisting of weak structural components.

Different components of EPS were correlated with various granular sludge properties in an anaerobic sulfidogenic granular sludge reactor [55]. The protein to polysaccharide ratio correlated positively with the loosely-bound EPS to tightly-bound EPS ratio. As the relative amount of proteins and loosely-bound EPS increased so did the hydrophobicity. However, the mean particle size decreased with increases in the relative amount of proteins and polysaccharides. Thus, an increase in the protein content of AGS particles may lead to enhanced microbial adhesion and more compact granule particles characterized by a dense EPS structure.
The role of individual components of the EPS matrix on AGS stability has still not been clarified. One of the reasons for this could be the extraction method. For instance, complete and representative extraction of EPS from AGS could be difficult due to the compact structure of granules [56]. Several extraction methods have been proposed including high-speed centrifugation, ultrasound-formamide-NaOH, or thermal, either individually or in combination. Due to the high variability of extraction methods, it has been suggested that there could also be great variability for the value of EPS obtained. Therefore, misrepresentation may occur if the role of EPS is assessed simply by these measured values.

*Ion exchange and biologically induced precipitation*

Initiating microbial adhesion is a critical step in attaining the compact structure of aerobic granules. Initiating microbial adhesion can be thermodynamically unfavourable in some cases since the electric repulsion of negatively charged, natural surfaces such as cells and certain EPS molecules may be greater than the attractive forces [57]. For this reason, anionic and cationic compounds must interact with one another to hold microorganisms and other EPS components together. Divalent cations (e.g., Ca\(^{2+}\), Mg\(^{2+}\)) help stabilize the microorganism-EPS complex via divalent cation binding [51]. On the other hand, monovalent cations (e.g., Na\(^+\), K\(^+\), NH\(_4^+\)) could exchange with divalent cations in the granular sludge complex, thereby replacing divalent cations and destabilizing the microorganism-EPS complex [58]. In general, monovalent to divalent cation ratios above 2 can lead to poor aggregation properties. However, compared to CAS and AAO-dominated granules, AGS particles had 4-9 times higher ammonium adsorption capabilities at an equilibrium ammonium concentration of 30 mg-N/L [59].
Other chemical exchange mechanisms, such as the exchange of potassium from potassium magnesium phosphate (i.e., K-struvite) in the granular sludge complex with ammonium in bulk solution, have been observed and contributed to the overall removal of nitrogen and phosphorus [60]. It has been suggested that small increases in the fraction of precipitates, as low as 1% to 5% of total granule volume, would significantly increase the density of aerobic granules and thereby their settling velocity [41]. Phosphorus precipitates, such as biologically induced hydroxyl-apatite (i.e., Ca$_5$(PO$_4$)$_3$(OH)), have been repeatedly detected in the core of aerobic granules [61], [62]. Other studies have observed elevated levels of other calcium precipitates, such as aragonite, and thereby suggested that elevated calcium levels in the wastewater may help induce granulation [63]–[65].

Components of the EPS matrix of aerobic granules have also induced the formation of struvite, another phosphorus precipitate commonly found in aerobic granules [60]. Alginate-like exopolysaccharides specifically, which are characteristic of aerobic granules, were found to induce the formation of potassium/ammonium magnesium phosphates (i.e., struvite). High concentrations of calcium precipitates specifically have been reported within the cores of granular sludge. It has been shown that concentrations of iron, magnesium, and aluminium remained constant in the granules during start-up whereas calcium increased [66], [67]. It was also observed that granules containing calcium precipitates had stronger structures and larger diameters, associated with a negative impact on bioactivity based on specific oxygen uptake rates [66], [68]. Larger granules, specifically greater than 0.45 mm, have also been shown to have slower denitrification rates than granules between 0.28-0.45 mm [69].
Microbial community and nutrient removal capabilities

Arguably, the most interesting aspect of aerobic granular sludge is the layering and interactions of different microorganisms throughout granule depth. All granules are characterized by a concentric multi-layered structure, which must contain channels and pores for the transport of oxygen and substrates [70]. With proper mass transfer conditions in place, aerobic, anoxic, and anaerobic zones characterized by a DO gradient across granule depth [71], [72]. This differentiation allows for simultaneous carbon, nitrogen, and phosphorus removal by a suite of microorganisms in one granule [73].

Typically, granules consist of an outer, 0.1-0.2 mm aerobic layer overtop anoxic/anaerobic layers. Aerobic microorganisms such as heterotrophs, AOB, NOB, PAO, and GAO grow in the outer layer while deeper layers contain facultative denitrifiers, denitrifying DPAO, and DGAO [50], [74]–[76]. Various other specialty microbes such as AAO, DAMOA, DAMOB, and methanogens have been observed within AGS [72], [77]. Nitrite/nitrate generated by AOB and NOB penetrates the anoxic layer, which acts as a nitrite/nitrate sink. The oxidized inorganic nitrogen is removed by conversion to nitrogen gas through denitrification, thereby creating a nutrient gradient where nitrite/nitrate travels from the aerobic to the anoxic layer. Denitrification and phosphorus removal pathways can be coupled through DPAO. It was observed that DGAO were the main microorganisms responsible for nitrate reduction in AGS reactors operated at 20°C and 30 °C [72]. It was also observed that a significant fraction of the nitrite was reduced to nitrogen gas via DPAO, specifically Clade II Candidatus accumulibacter phosphatis.
If AAO develop within the anoxic layer, nitrite can also be utilized to oxidize residual ammonium to nitrogen gas with approximately 10% residual nitrate [25]. The nitrate produced by AAO can be utilized by any of the facultative microorganisms to oxidize organic matter to carbon dioxide. Encouraging establishment of AAO within aerobic granules has been characterized by higher ammonium effluent concentrations (e.g., >1 mg/L) and low (e.g., 0.5 g-COD/g-N) carbon to nitrogen ratios [77], [78]. Furthermore, the ratio of NOB to AOB in AGS systems has been observed to be higher than in CAS systems, which does not benefit AAO establishment since they compete directly with NOB for nitrite [79]. These results demonstrated that there is complex nutrient cycling within AGS particles, where some NOB are either growing mixotrophically using organics for nitrate reduction or denitrifiers are reducing nitrate to nitrite within AGS particles and thereby supplying additional nitrite for NOB without the additional ammonium for AOB.

The diverse microbial community of aerobic granular sludge, coupled with its ability to generate nutrient gradients, allows it to perform complete nutrient removal in a single tank simply by switching airflow on or off. Completely aerobic granular sludge processes, those that do not implement an anaerobic influent distribution phase, have demonstrated excellent organic carbon and nitrogen removal capabilities. Processes that do have an anaerobic contact phase, however, have also exhibited EBPR. Thus, AGS becomes an ideal process for removal of carbon, nitrogen, and phosphorus from wastewaters. Furthermore, the slow growth of microorganisms in stable granular sludge systems as well as the constant recycling of organic and inorganic compounds for oxidation and reduction processes throughout the granule contribute to lower sludge yield and reduced dependency on external carbon and/or metal salt addition to meet stringent nutrient limits compared to conventional activated sludge systems.
Dead cells and precipitates can also exist under anaerobic conditions near the core of aerobic granules [80]. Being an active microbial community, AGS is constantly growing and therefore increasing in diameter, unless mechanically broken apart first. As granular sludge diameter increases the extent to which oxygen, nitrite, and nitrate penetrate decreases, resulting in the development of anaerobic conditions in the interior of granular sludge particles [72]. Depending on the sludge age of the anaerobic core, it is possible that microorganisms capable of degrading EPS in the granular sludge matrix become established. It has yet to be demonstrated whether this is a common mechanism for self-maintenance of individual granular sludge particles or a mechanism that induces instability, ultimately leading to the breakdown of larger granular sludge particles.

Since the aerobic layer does not usually exceed more than 0.2 mm, the diversity of the microbial community should increase as average granule diameter surpasses 0.4 mm. Furthermore, as bulk DO concentrations decrease, the aerobic layer is expected to decrease in size, thereby leading to more diverse microbial communities in smaller granules (i.e., <0.4 mm). Microbial diversity as well as community structure would directly influence the performance of aerobic granules. For instance, analysis of full-scale granular biomass indicated that PAO had a higher density than GAO, leading to significantly higher settling velocities for PAO dominated granules [41]. Thus, out-selection of GAO could improve AGS performance. Furthermore, the kinetic rates and stoichiometry of individual metabolic pathways would govern granule stability.
The higher the microbial community yield, the more quickly the control volume (i.e., the granule volume) would be “replaced” by new biomass. Higher microbial community yields would therefore make granules more susceptible to mass transfer limitations due to excessive growth of microorganisms and their byproducts, such as EPS [11], [43]. Thus, continuously aerated AGS systems at high DO would be less stable than AGS systems with anaerobic influent distribution operated at low DO. The elimination of aerobic/anoxic substrate utilization by heterotrophic microorganisms would be desired, while anoxic utilization of substrate by PAO and GAO would be preferred.

It was demonstrated, on a molecular level, that AGS communities are much more efficient than flocculent communities at utilizing nutrients [81]. Based on the number of significantly enriched proteins assigned to Clusters of Orthologous Groups, it was shown that the AGS metagenome had significantly higher transport and metabolism related proteins (i.e., for carbohydrates, amino-acids, coenzymes, lipids, and inorganic ions) while flocculent communities had more proteins related to translation, ribosomal structure, biogenesis, energy production, and conversion. In general, this means that the flocculent community was actively growing without expending too much energy on substrate utilization while the AGS community had to invest energy into adenosine triphosphate consuming transport mechanisms for substrates with low energy return [82].
**Surface layer of granular sludge**

Contrary to common depictions, AGS is not always made up of uniform spherical particles with smooth surfaces. The structure of granular sludge, including the microorganisms in the outer surface layer, is dependent on both wastewater characteristics and operational strategies. For instance, uneven granule surfaces can develop when AGS is used to treat wastewater with high slowly-biodegradable, particulate organic loads [44]. This is because slowly-biodegradable particulates sorb to the surface of granules and gradually release bioavailable organic matter via hydrolysis. The hydrolyzed organic matter is then used within the vicinity of the original particulate organic matter that had sorbed to the granule, resulting in a microcosm of heterotrophic activity. At steady-state an equilibrium would be obtained between new slowly-biodegradable, particulate matter attaching to the surface of granular sludge and older particulate matter being utilized by the aerobic heterotrophs surrounding it.

When the load of slowly-biodegradable, particulate matter is low, there would be no significant effect on granule surface structure and the outer layer may appear smooth and compact, if other operating parameters are within bounds. As the quantity of slowly-biodegradable, particulate matter increases, however, the rate of degradation by aerobic heterotrophs will be lower than the rate of attachment, resulting in the retention of particulate organic matter to the outer surface and the appearance of an uneven granular structure [83]. The excessive substrate sorbed to granule surfaces may also encourage the development of filamentous microorganisms feeding off the aerobically hydrolyzed substrate. Furthermore, the presence of septic wastewater could also encourage the growth of filamentous microorganisms on granule surfaces [43], [84]. The presence of filamentous microorganisms would increase the drag force of AGS and decrease settling velocity of granules, resulting in poor-quality effluent.
Factors affecting granule formation and stability

*Alternating “feast” and “famine” conditions*

Biomass with high internal substrate storage response, such as PAO and GAO, are thought to benefit from alternating conditions of excess substrate (i.e., feast phase) and lack of substrate (i.e., famine phase) – Figure 1. It has been suggested that this is because rapidly growing heterotrophic microorganisms experience higher decay under extended aerobic conditions, or the famine phase. Thus, growth of microorganisms capable of PHA storage during the feast phase would be favoured, especially since PHA storing microorganisms usually have lower decay rates. It is unlikely that PHA storing microorganisms would be selected for in a continuously aerated AGS-SBR.

*Figure 1 Depiction of feast and famine phases by OUR profiles in a continuously aerated CAS-SBR (left) and continuously aerated AGS-SBR (right) with and without extended feast conditions due to high pCOD in the influent. The endogenous respiration phase in OUR profiles can be used to monitor the development and duration of the famine phase.*
Simulations using BioWin\textsuperscript{TM} software of a completely aerobic plug-flow system (i.e., five equally sized bioreactors in series) demonstrated that the ratio of PAO to OHO would be $<<1\%$ – Figure 2 [86]. Rather, selection of PHA storing microorganisms would depend on maximizing the availability of VFA to PAO and GAO under anaerobic conditions. It should be noted that wastewaters deficient in nitrogen, phosphorus, or other elements for cellular growth could also encourage PHA storage [85]. In municipal wastewater, there is typically not enough bCOD to fully remove TN or TP. If the ratio of bCOD to TN to TP is above 100:5:1 there should be no nitrogen or phosphorus deficiency. However, if AGS was applied to wastewaters with excess COD and low TN and TP, such as brewery wastewaters, PHA may be encouraged under completely aerobic conditions [11].

Figure 2 Estimated PAO to OHO ratio for a modelled biological treatment facility with increasing SRT and various configurations at a 12 h HRT. All models had typical primary influent characteristics of 500 mg-COD/L, 40 mg-TN/L, and 10 mg-TP/L as input. Fermentation was modelled at a 3 d SRT receiving primary sludge from a clarifier removing 70% of influent TSS.
The development of a famine phase depends on wastewater characteristics and OLR [87]. For instance, the presence of slowly-biodegradable, particulate matter may extend the duration of the feast phase by supplying a constant source of readily biodegradable organics. Furthermore, treatment plants that are nearing design capacity may also lack the required quantity of biomass and contact time required to rapidly reduce the concentration of bioavailable substrate, leading to shorter starvation periods and potentially higher process instability. An imbalance in the feast and famine duration, with prolonged feast phase at the expense of a short famine phase, led to granule disintegration in an AGS-SBR with anaerobic influent distribution [88]. To maintain stable AGS, it was suggested that the feast period should not exceed 25% of cycle length.

**Hydrodynamic shear forces**

Hydrodynamic shear forces may be defined as forces generated due to the friction between the bulk liquid and granules surface. The hydrodynamic shear force has typically been quantified by USAV in lab-scale systems. This parameter is calculated as the airflow divided by the cross sectional area of the aerated zone [43]. The exact value for USAV varies in the literature. For instance, some did not observe granulation when USAV were less than 1.2 cm/s [89]. Others found that unstable granules developed when USAV was less than 2.4 cm/s [90]. It has also been observed that AGS was not stable at USAV below 1 cm/s [91]. The authors stated that low aeration rates resulted in oxygen limitation during aeration, thereby increasing the duration of the feast period and shortening the famine period. However, most of the studies linking hydrodynamic shear force to granulation potential were completed in continuously aerated AGS systems.
More recently, it was demonstrated that high hydrodynamic shear forces are not always a necessity for granulation, although they may benefit AGS stability in some cases [43]. Specifically, high shear may remove fast growing heterotrophic microorganisms, while also preventing the accumulation of particulates originating from the influent or of microbial origin, on granule surfaces. At higher OLR the potential for fast-growing, heterotrophic microorganisms to grow in bulk solution or on the surface of granules is higher. Thus, at higher OLR, higher hydrodynamic shear forces would displace detrimental matter from granule surfaces and prevent the formation of a loose outer structure that could lead to poor settling properties and therefore AGS instability. At lower OLR, successful granulation has been observed at USAV below 0.5 cm/s [43].

Influent distribution

In general, AGS-SBR systems may be categorized into one of two types of influent distribution: 1) continuously-aerated; and 2) anaerobic influent distribution. The method of influent distribution would influence the morphology and microbial community of aerobic granules. It was observed that a long feeding time, up to 50 min, through a settled bed of granules was favourable over a short feeding time during mixing [92]. The two most important reasons were that the application of a long, influent distribution period at large-scale treatment plants simplified SBR scheduling, while also selecting for PAO and GAO due to the alternating aerobic and anaerobic phases. Granulation from CAS inoculum was successful under a 20 min anaerobic-40 min aerobic split feed strategy, although the granules exhibited poor phosphorus removal [93]. It is generally agreed that the basic principle of stable granulation is the selection of slower growing bacteria, and distributing substrates throughout the granule during anaerobic influent distribution encourages the selection of slow-growing microorganisms [94].
In the case of wastewater rich in easily biodegradable soluble substrates, when influent was distributed anaerobically with a plug-flow regime, the substrates are taken up by PAO or GAO and converted to internal storage polymers, such as PHA, that are used for growth at a relatively slow rate in subsequent aerobic/anoxic phases [92]. At the beginning of the cycle, PAO and GAO near the granule surface take up easily biodegradable substrates. As anaerobic feeding continues, the PAO and GAO near the surface become saturated with internally stored polymer and the easily biodegradable substrates penetrate further into the granule. Thus, substrate reaches PAO and GAO deeper within the granule. This leads to uniform biomass production throughout the granule, thereby stabilizing the granule structure. Since oxygen will not penetrate into the deeper regions of granules, even during vigorous aeration, storage polymers can be oxidized by other electron acceptors such as nitrate, ensuring optimal phosphate and nitrogen removal [95].

Conversely, an aerated influent distribution strategy could lead to diffusion limitations, commonly characterized by a surface layer of filamentous microorganisms. During the feast phase, when bulk liquid substrate concentration is high, the substrate would be rapidly utilized for growth limited to the outer aerobic layer of the granule [96]. Furthermore, the rapid, aerobic consumption of septic substrates in the presence of oxygen is well known to encourage the formation of filamentous microorganisms, which could foul the surface of granules [84]. Filamentous growth will have detrimental effects on the settling properties of AGS, and thus negatively impact effluent quality [94], [97]. In these cases, increased hydrodynamic shear may be required to ensure smooth and stable granulation. Significant surface fouling, due to the proliferation of filamentous microorganisms, was observed when organic loads exceeded the capacity for anaerobic uptake, even with pure acetate as substrate [43]. Overall, granules formed under an aerobic influent distribution are more prone to degranulation [96].
Particulate substrates, often present in high quantities in industrial wastewaters, also impact AGS formation and stability. Once granules have developed, particulate substrates can be hydrolysed at the granule surface [83]. The porous structure of granular sludge acts as a filter, retaining particles larger than the pore at the granule surface. Hydrolysis products created during anaerobic influent distribution will be converted into storage polymers, thereby selecting for PAO and GAO. However, depending on the anaerobic hydrolysis rate as well as the quantity of particulate matter, aerobic hydrolysis will also occur. Aerobic hydrolysis will supply a steady source of biodegradable matter and extend the duration of the feast phase. Aerobic hydrolysis product will then be used for growth by heterotrophic organisms at the surface of granules, causing limitations in substrate diffusion [98]. Limitations in substrate diffusion could induce filamentous outgrowth, result in less stable AGS, or produce higher suspended solids in the effluent. In this respect, it was noted that extended famine regimes, by decreasing OLR, could enhance AGS stability [45], [88].

Lastly, if easily biodegradable substrates are slowly supplied to a mixed aerobic environment, severe limitations in substrate diffusion may develop and filamentous microorganisms could develop. That is because high, biodegradable substrate availability in bulk solution during aeration is favourable for the growth of fast-growing, heterotrophic microorganisms. Filamentous growth will have a detrimental impact on the settling properties of AGS, and thus negatively impact effluent quality [94], [97]. Thus, maximizing anaerobic utilization of biodegradable substrate prevents the proliferation of fast-growing, aerobic heterotrophs and encourages uniform granules with smooth and compact surfaces.
Selective wasting

In most AGS studies the selection pressure was related to the settling time. Several studies have demonstrated that shorter settling times enhance aerobic granulation [99], [100]. A higher initial washout of biomass induces the production of excess EPS, secreted by functional strains, without a strong challenge for substrate by competing strains that have been washed out. However, some studies have suggested that very short settling times lead to premature washout of certain non-flocculated strains, thereby yielding low biodiversity of granules [101]. It was also observed that too rapid a decrease in settling time leads to washout of desired bacterial guilds such as PAO, AOB, and NOB [102]. It has been suggested, as an option to avoid excessive biomass loss during start-up, that the settling time be linked directly to the observed washout, as well as to recirculate sludge from a secondary clarifier in case of a massive washout [103].

Although short settling times cause the washout of poorly-settling, free-living microorganisms while retaining fast settling particles, short settling times discharge the selectively wasted light fraction of biomass with the effluent [104]. This can result in high effluent solids concentrations, more than discharge limits. A better method of granulation would be to employ a discrete selective wasting regime apart from effluent discharge [105]–[107]. For instance, it was demonstrated via FISH that more dense granules located at the bottom of the settled sludge bed contained considerably more PAO compared to the lighter granules on top which were dominated by GAO [107]. By selectively wasting biomass exclusively from the top of the settled sludge bed TP removal efficiency increased from 71% to near 100%. By separating selective wasting from the effluent discharge, treated wastewater solids can be less than 5 mg/L and higher removal efficiencies can be obtained [18].
Organic loading rate

OLR is defined as the ratio between organic substrate load, calculated as the product of flow and substrate concentration, and volume of the reactor receiving the substrate. The F/M takes the volumetric representation of OLR and converts it to the ratio of organic substrate load and “active” biomass, usually as MLVSS. In continuously aerated AGS systems, where the entire cycle, including the influent distribution, is under aerated conditions, OLR and F/M seemed to be important for aerobic granulation. In one instance it was determined that the optimum OLR for granule formation was around 2.5 kg-COD/(m³•d) [108]. In another case, the formation of AGS in SBRs supplied glucose-based substrate determined that higher F/M, up to 1.1 kg-COD/(kg-VSS•d), promoted faster formation of granules with larger average diameter [109]. Similarly, another study analyzed the granulation process at OLRs between 1.5-4.5 kg-COD/(m³•d) [110]. It was observed that higher OLR resulted in faster formation of larger and looser granules, while a lower loading rate resulted in slower formation of smaller and more dense granules. It was even observed that granulation could not be achieved at an OLR lower than 2 kg-COD/(m³•d) for continuously-aerated SBRs [89]. In contrast, another study successfully cultivated granules at an OLR of 1-1.7 kg-COD/(m³•d) in a continuously-aerated SBR, however almost one year was needed to obtain mature granules [111].

The role of OLR and F/M appeared to be reduced with the introduction of an anaerobic influent distribution phase. With anaerobic influent distribution, it was demonstrated that stable AGS has been generated at lower OLR than continuously-aerated SBRs. Full-scale Nereda® reactors operate with OLR ranging from 0.65-0.85 kg-COD/(m³•d) [31]. Similarly, stable AGS was cultivated on proteinaceous substrate within a month at an OLR of 1.4 kg-COD/(m³•d) [43]. Stable AGS was also cultivated within three weeks at an OLR of 1.4 kg-COD/m³•d while
treating brewery wastewater [88]. Based on the reported studies, it can be concluded that OLR and F/M are not crucial parameters for aerobic granulation. However, the impact of high OLRs and F/M is more significant in continuously-aerated AGS systems. These results clearly indicate that there are more important factors influencing aerobic granulation than the OLR.

*Other environmental factors*

The main environmental factors affecting biological processes, including AGS, are DO, ORP, pH, and temperature. So far, the literature has suggested that DO was not a dominating factor for granulation despite a single claim that DO, instead of hydrodynamic shear stress, governed the granulation process [112]. Instead, DO was suggested to have an impact on the layering and stratification of microbial communities. Since an oxygen gradient develops across the depth of granular sludge particles, anaerobic, anoxic, as well as aerobic layers could coexist within a single granule at any given time.

Stratification clearly impacts nutrient removal processes due to the simultaneous presence of varying ORP conditions and therefore the capacity for SND. The aerobic layer of granules was demonstrated to be strictly related to bulk DO concentrations. It was demonstrated, for granules of equal diameter, that oxygen penetrated further into the granule when bulk DO concentrations were higher. Thus, if DO in the bulk liquid was too high the development of anoxic or anaerobic layers could be inhibited, especially for smaller granules. Similarly, it was observed that reduction of DO to 40% saturation during the aeration phase significantly improved TN removal efficiency [95].
Some authors found that acidic and neutral solutions in AGS reactors had a significant impact on granulation [113]. Concerning pH, it was observed that fungi dominated aerobic granules at a pH of 4, and that the granules were as large as 7 mm. Fungi resulted in faster granulation by providing a filamentous matrix for the microbial community to grow upon. However, if the fungal growth was not controlled, their irregular patterns could lead to system instability. At pH higher than 8, granules were dominated by bacteria and granule size only reached 4.8 mm. Neutral medium and bacterial dominance resulted in granules that had a more compact structure. Other studies have observed that alkaline pH induced granule disintegration and resulted in increased effluent solids, as well as a significant decrease in the size of AGS [114]. The same study suggested that acidic pH did not have a significant impact on AGS stability. Changes in chemical structure and composition of the EPS matrix were suggested as the main factors inducing granule instability under high pH.

Temperature, in addition to affecting bacterial growth kinetics, also influences the settling velocity because of changes in the viscosity of bulk solution. It was observed that, in an SBR operated at 8 °C, granules with irregular shape and excessive filamentous bacteria developed which caused unstable operation by severe biomass washout [87]. The authors concluded that aerobic granulation at low temperatures was not practically feasible. In general, microbial processes are slower at lower temperatures, and the slower rates could have limited granulation at lower temperature [115]. Another study demonstrated that granule settling velocities are also significantly reduced at lower temperatures [40]. On the other hand, operating at high temperature (e.g., >30 °C) could also inhibit granulation by preventing agglomeration of EPS and increasing the solubility of chemical species (e.g., iron, magnesium, phosphorus) that have been detected in the core of AGS [116].
Design considerations and control strategies

Aerobic granulation can occur in a wide range of scenarios with different operating parameters. However, certain variables do seem to be more significant for aerobic granulation and long-term granule stability. Based on the literature, several operating parameters were compiled according to their importance for granulation – Table 5. Among these, anaerobic influent distribution was almost unanimously considered essential for the formation of stable AGS [92], [94]. Duration of the anaerobic influent distribution phase will depend on the OLR and wastewater characteristics. Anaerobic influent distribution should be designed to maximize the conversion of organic matter in the influent to intracellular polymeric substances, such as PHA. Fifty minutes to one hour is typically enough for the anaerobic influent distribution phase when treating municipal wastewater.

Table 5 Summary of the main operating conditions affecting aerobic granulation.

<table>
<thead>
<tr>
<th>Operating parameter and other factors</th>
<th>Impact on the process</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaerobic influent distribution</td>
<td>✓✓✓✓</td>
<td>Improves AGS stability and allows for nutrient removal.</td>
</tr>
<tr>
<td>Selective wasting</td>
<td>✓✓✓</td>
<td>Retains AGS, washes-out undesirable biomass.</td>
</tr>
<tr>
<td>Famine conditions</td>
<td>✓✓</td>
<td>Allows for EPS modification, washes-out biomass with high decay rates.</td>
</tr>
<tr>
<td>Feast conditions</td>
<td>✓✓</td>
<td>Diverts carbon from growth to EPS production.</td>
</tr>
<tr>
<td>Hydraulic shear forces</td>
<td>✓</td>
<td>Stabilizes AGS at high OLR.</td>
</tr>
<tr>
<td>SRT</td>
<td>✓</td>
<td>Prevents excessive aging of granules, stabilizes AGS at high OLR.</td>
</tr>
<tr>
<td>DO</td>
<td>✓</td>
<td>Not too low to prevent the core weakening, not too high to favour a proper layered structure.</td>
</tr>
<tr>
<td>Organic loading rate</td>
<td>✓</td>
<td>Higher OLRs can accelerate start-up.</td>
</tr>
<tr>
<td>pH</td>
<td>✓</td>
<td>Alkaline pH shock may destabilize AGS.</td>
</tr>
<tr>
<td>Temperature</td>
<td>✓</td>
<td>Low temperature delays the granulation process.</td>
</tr>
<tr>
<td>Substrate composition</td>
<td>×</td>
<td>AGS developed with several organic substrates.</td>
</tr>
</tbody>
</table>

Legend: ✓✓✓✓ = Fundamental; ✓✓✓ = very important; ✓✓ = important; ✓ = not crucial; × = not relevant.
It was observed that high OLR encouraged proliferation of filamentous microorganisms, initially favouring a rapid increase in the size of the granules due to the ability to use filaments as structural components [50], [117]. The filamentous microorganisms, however, also resulted in granules with loose structures and therefore lower settling velocities. In this respect, it was suggested that the selection of slow-growing organisms (e.g., PAO and GAO) could enhance granule stability [92]. The same authors suggested that a long feeding time through a settled sludge bed appeared feasible and was favourable over a short feeding time.

It has been hypothesized that a certain “critical” granule size will be achieved at steady-state under certain operational conditions [118]. Due to the consequences of particle collision and structural weaknesses, granules larger than the critical size are expected to break apart until reduced to less than or equal to the critical size. Conversely, the growth of small granules in steady-state AGS reactors equals the rate at which large granules are being reduced by breakage or attrition. This cycle of granulation and degranulation becomes a dynamic process that is constantly evolving. Similarly, it was demonstrated that a SRT of granular sludge longer than 14 days leads to granule deterioration [119]. To mitigate the disintegration of older granules, it was suggested that selective discharge of old, and therefore large, granules from the bottom of the reactors (e.g., wasting 5% of the settled bed daily) be performed [120].

The reason for larger granules weakening is partly due to mass transfer limitations leading to a weakened inner core of the granule. It had been suggested that mass transfer limitations of large granules could produce an anaerobic core and thereby stimulate the activities of anaerobic strains, hence breaking apart large granules [121]. One study claimed that the breaking of mature granules was due to the clogging of pores and channels in the granules, hence hindering nutrient intake by the microorganisms [122]. In this context, it has been demonstrated
that excessive EPS production contributed to the clogging of pore space in the granules, causing their breakage in the long run [45]. An extended famine period encouraged greater EPS consumption by bacteria, thus limiting the clogging of pores, and allowed for the long-term operation of stable aerobic granules. It has also been noted that granule disintegration at very high OLR (i.e., >21.3 kg-COD/(m³•d)) is the result of microorganisms losing their capacity to produce proteins or polysaccharides at the applied OLR [117].

Particulate organic matter and other slowly-biodegradable organic substrates can offset the benefits of anaerobic influent distribution. Due to slow anaerobic hydrolysis rates, slowly-biodegradable substrates may not be fully converted to PHA during anaerobic influent distribution. Therefore, the slowly-biodegradable substrates would contribute to a continuous supply of bioavailable substrate via aerobic hydrolysis during aeration [83]. The growth of aerobic heterotrophs would thereby be promoted in a confined location since particulate and colloidal substrates would be concentrated in the bulk solution or near the granule surface. Overgrowth at the granule surface could lead to mass transfer limitations and the loss of nutrient removal [43].

Two approaches have been suggested to prevent the proliferation of microorganisms on the surface of granules that feed off aerobically hydrolyzed substrate. The first approach is to apply an extended famine phase by increasing the duration of the aerobic period. The feast phase, the period of time in which hydrolysis is occurring and substrate is available near the granule surface, should not be more than 20% of the aerobic period [88], [123]. The second approach would be to increase the applied hydrodynamic shear force in order to detach filamentous overgrowth at the granule surface [43]. From a design point of view, the first
approach implies an increase in capital cost due to larger reactor volumes while the second implies an increase in operating costs for electricity to power blowers or mixers.

The last crucial parameter for aerobic granulation is selective wasting. It is considered a fundamental parameter because it allows for the out-selection of slowly settling flocculent biomass and the retention of rapidly settling AGS. Typically, a critical settling velocity is applied such that only particles with higher settling velocities will be retained within the reactor. The settling velocity applied for selective wasting can be as high as 8 cm/min whereas conventional clarification following BNR activated sludge are designed with settling velocities of 1.7-2.2 cm/min [10].

Application of AGS to municipal wastewater

*Municipal wastewater characteristics*

Municipal wastewater is made up of both soluble and particulate, organic and inorganic matter. Proteins, carbohydrates, lipids, and fibers contribute to the particulate organic fraction, while readily available soluble organic matter is comprised of VFA, alcohols, and other simple carbohydrates [10]. Depending on the configuration of sewer systems municipal wastewater can contain low or high quantities of bioavailable substrates. Larger sewersheds located on relatively flat terrain in warmer climates will generally yield septic wastewater with lower biodegradable fractions, having already been utilized within the sewer system [5]. On the other hand, municipalities in mountainous regions with steep terrain and lower temperatures will provide treatment facilities with fresh, unaltered wastewater.
The fraction of soluble biodegradable substrate in the wastewater is particularly important for AGS processes. The higher the fraction of soluble biodegradable substrate, the more likely the AGS process is to be stable. This is because fresh wastewater can be introduced into an anaerobic contact phase where readily bioavailable substrate is taken up and converted to internal storage product by accumulating bacteria. Thus, there will be less residual organic matter in bulk solution passing through to aerobic conditions if more of the substrate is soluble biodegradable. By diverting a majority of the initial organic matter to internal storage products, it is less likely that a fast-growing, aerobic heterotrophic culture will develop on granule surfaces, resulting in mass transfer limitations and process instability [43], [92].

Furthermore, municipal wastewater can be classified as low-, medium-, or high-strength depending on how water conservative municipalities are, as well as how much inflow and infiltration is experienced. Municipalities that use less water and have separate sewers tend to have stronger wastewater that contains higher concentrations of organic and inorganic matter. It has been speculated that AGS, which derives benefits from the development of nutrient gradients across granule depth, performs better on high-strength wastewaters than low-strength, diluted wastewaters [120]. Higher concentrations facilitate diffusion and therefore promote more rapid uptake and utilization of nutrients by aerobic granular sludge.
Optional and required pre-treatment of municipal wastewaters

The extent of municipal wastewater pre-treatment is still largely debated in literature and practice. Personal communication with leading environmental consultants has yielded contradictory responses. Some swear by removing as much solid content as possible up front through primary treatment while others have argued that AGS only works on primary influent. These claims are based on individual experiences with demonstration facilities likely confounded by site-specific variables that have not been accounted for in design and operation of the demonstration facilities. Full-scale Nereda® installations have worked successfully for many years on primary influent, demonstrating that the AGS-SBR process is a robust technology capable of handling the additional particulate loads and variability associated with primary influent [14], [31].

It is expected that the only necessary pre-treatment before any AGS process would be screening and grit removal. Due to the nature of AGS processes, which select dense particles, the breakthrough of grit from pre-treatment could result in significant accumulation of undesirable inert material. Thus, it is very important that good grit removal with enough redundancy be designed for AGS processes. Optional pre-treatment for AGS processes includes the removal of particulate organics from the wastewater. A summary of the pre-treatment options and their applications is provided in Table 6.
Table 6 Summary of AGS-SBR pre-treatment configurations based on municipal wastewater characteristics.

<table>
<thead>
<tr>
<th>Pre-treatment</th>
<th>Process diagram</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screening + Grit removal</td>
<td><img src="image1" alt="Diagram" /></td>
<td>For municipal wastewater with TSS and VSS capable of adsorption and degradation by AGS without disrupting process stability.</td>
</tr>
<tr>
<td>Screening + Grit removal + Sedimentation</td>
<td><img src="image2" alt="Diagram" /></td>
<td>For municipal wastewater with TSS and VSS exceeding the capacity of AGS, having adequate residual COD to TN ratio for nutrient removal.</td>
</tr>
<tr>
<td>Screening + Grit removal + Sedimentation + Fermentation</td>
<td><img src="image3" alt="Diagram" /></td>
<td>For municipal wastewater with TSS and VSS exceeding the capacity of AGS, as well as COD deficiency for nutrient removal.</td>
</tr>
</tbody>
</table>

The removal of additional influent solids, besides grit, before aerobic granular sludge reactors has many plausible arguments both for and against it. Reasons for good solids removal up front include the separation of slowly-biodegradable matter, which can lead to uneven surface structure, and the removal of dense, inert particles that could accumulate in granular sludge reactors. Diversion of settleable organics away from mainstream processes has been practiced for increased mainstream capacity and beneficial reuse in the side-stream [5]. On the other hand, it has been argued that higher wastewater solids content can help initiate, or is even necessary for, granulation. High density, unbiodegradable inert particles found in primary influent, which mostly settle out during primary treatment (i.e., 50-70% removal), may contribute to more rapidly settling granular sludge particles if the inert particles become associated with the granular sludge matrix.

36
It has also been suggested that allowing more slowly biodegradable particulate organic matter through to the granular sludge process can alleviate problems associated with low-strength wastewaters. Nevertheless, with properly designed cycles that account for the peak organic load in the design year, AGS-SBRs can adequately treat primary influent with particulate organic substrates to exceptional effluent quality [45], [124]. Thus, the application of properly balanced feast and famine phases allows for additional organics in the primary influent to be beneficially used for nutrient removal without the particulate load negatively influencing the AGS process.

Generally, particulate substrates are hydrolysed at the surface of granules during steady state in properly designed AGS-SBRs. Municipal wastewater with abnormally high particulate substrate can negatively impact AGS stability [83]. The porous structure of granular sludge acts as a filter, retaining particles larger than the pore size at the granule surface. Hydrolysis products during anaerobic influent distribution would be converted into storage polymers, thereby selecting for PAO and GAO. However, depending on anaerobic hydrolysis rates as well as the quantity of particulate matter, aerobic hydrolysis will also occur. Aerobic hydrolysis would supply a steady stream of biodegradable matter that would be used by heterotrophic organisms at the surface of granules [43], [98]. This has induced filamentous outgrowth, resulted in less stable granules, and generated higher suspended solids in the liquid phase. If wastewater characterization determined that influent wastewater was high in particulate matter, primary treatment should be considered.
Primary treatment could include sedimentation, micro-screening, or CEPT. The removal of organic matter may result in low carbon to nitrogen ratios, and therefore poor nutrient removal even with SND. If the wastewater was determined to be high in particulate matter, as well as carbon deficient after primary treatment, fermentation of primary solids could be used to increase the amount of readily available substrate sent to the AGS process [125]. Fermentation products, such as acetate and propionate, are easily utilized by AGS and contribute to their stability [126], [127].

*Operational considerations for municipal wastewater treatment*

An efficient, stable AGS-SBR process treating municipal wastewater must consider a few design factors. The following discussion will include operational considerations for municipal wastewater treatment with AGS as it pertains to: 1) cycle configuration; 2) OLR; 3) selective wasting; and 4) anaerobic influent distribution. Cycle timing must be designed to provide adequate oxidation of soluble and particulate organic compounds within diurnal flow and load variation. If higher loads of organics exceed the anaerobic uptake capacity of biomass during a single cycle, the residual organics in bulk solution would pass on to the aerated react phase and/or subsequent cycles. Under these conditions residual organics could promote the growth of fast-growing, aerobic heterotrophs, ultimately destabilizing the process if they are allowed to proliferate further [94]. Thus, AGS cycles should be designed with flexibility in order to address variable influent loads.
Depending on the degree of flow, load, and temperature variation observed, different design considerations can be made. These could include equalization and flow balancing, DO control strategies to ensure adequate “famine” conditions develop at higher observed loads, multiple draw points for selective wasting by a pump with variable frequency drive (i.e., remove more or less biomass considered to be part of the light fraction and therefore not desirable), and swing-capabilities allowing portions of the anaerobic influent distribution phase to be aerated [93], [124].

The literature varies in terms of recommended OLRs. For instance, the reported range on municipal wastewater has been from 0.6-1.4 kg-COD/(m³•d) [31], [43]. Most of the reported data from full-scale Nereda® facilities may not represent the design OLR, since most of their installations are relatively recent (i.e., first installation in 2012). Rather than a set OLR, AGS processes for municipal wastewater treatment should be designed and operated based on F/M. The flow and load of wastewater will continue to increase for municipalities experiencing population growth. Thus, the OLR will always be increasing. For this reason, operators of AGS facilities must understand the optimal biomass concentrations, relative to the incoming load, that will provide adequate nutrient removal and energy efficiency. In general, F/M ratios from 0.3-1.1 kg-COD/(kg-VSS•d) have been adequate for AGS operation [109]. However, operating above 1 kg-COD/(kg-VSS•d) may not be suitable as the risk for hydrous bulking increases.
Another important consideration includes selective wasting and the decoupling of granular solids retention time from free-living microorganisms. Ideally, the SRT of free-living microorganisms would be designed to be less than their minimum SRT. By doing so, the proliferation of fast-growing heterotrophs would ideally never occur. Given the inherent variability and characteristics of municipal wastewater, some substrate will always be available under aerobic conditions, thereby promoting the growth of free-living microorganisms and the development of a persistent light fraction of biomass. For this reason, selective wasting is important in order to manage, rather than completely out-select, the light fraction of biomass [128]. Designers and operators should consider that the more aggressive the selective wasting strategy (i.e., lower applied SRT), the higher the yield of the light fraction of biomass. Thus, selective wasting should be optimized to aid in the diversion of substrate to anaerobic utilization by microorganisms within AGS particles. The only way to decrease the yield of biomass, and therefore improve the performance of AGS, is to divert substrate away from direct aerobic utilization and towards anaerobic uptake [43], [94].

To effectively limit the growth of fast-growing, aerobic heterotrophs, AGS-SBRs should be designed to divert as much carbon as possible to internal storage product during the anaerobic influent distribution phase. In general, an anaerobic contact time of 50 min has been suggested as sufficient for anaerobic substrate conversion while treating municipal wastewaters [92]. However, more dilute wastewaters in areas with higher degrees of inflow, infiltration, and wet weather may require less anaerobic contact time [93]. Arguably, one of the most important factors is the configuration of influent distributors in AGS-SBRs with anaerobic influent distribution. Without proper hydraulic design, short-circuiting could occur and thereby disrupt optimal conditions for anaerobic uptake of substrate in the settled sludge bed.
Short-circuiting is detrimental since anaerobic diversion of organic carbon to internal storage product is desired [43], [129]. It is well known that the breakthrough of organic substrate in bulk solution to aerobic conditions negatively affects AGS stability in SBR. This could be due to many reasons, including the overgrowth of filamentous organism on the surface of granules or an accumulation of particulate matter. Both cases lead to mass transfer limitations and therefore increase the potential for granule disintegration. For municipal wastewater treatment, where stringent discharge limits must be achieved (e.g., TSS <5 mg/L, BOD₅ <5 mg/L, TN <5 mg/L, and TP < 0.5 mg/L), an AGS-SBR configuration with anaerobic influent distribution would be preferred – Table 7.

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Process diagram</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuously aerated</td>
<td></td>
<td>Influent is supplied over a very short period (i.e., &lt;5 min) during aeration. The settling time is also short (i.e., &lt;10 min) and selectively wasted biomass is discharged with the effluent. Feed and discharge occur separately.</td>
</tr>
<tr>
<td>Anaerobic influent distribution</td>
<td></td>
<td>Influent is supplied gradually over an extended period (i.e., 50 min) under anaerobic conditions. Discharge occurs with feed. After influent distribution feed, the selectively wasted biomass is removed, separating the light solids from the effluent.</td>
</tr>
</tbody>
</table>
Application of AGS to industrial wastewaters

Agro-food wastewater

Agro-food industries generate large volumes of wastewater from both the production line and cleaning practices. Some of these industries generate high salinity wastewater (i.e., fish-canning), wastewater with high suspended solids, as well as wastewater with high organic substrate concentrations. High-throughput treatment system are therefore required to comply with discharge limits (e.g., into the municipal sewer or direct discharge to surface water) or to obtain an effluent that is suitable for reuse within the industry or other nearby sites.

Brewery wastewater

Wastewater produced by breweries are characterized by high concentrations of bCOD derived from organic components (e.g., yeast, sugars, soluble starch, etc.). Nitrogen and phosphorus concentrations are typically dependent on the amount of yeast in the effluent [130]. Thus, brewery wastewater can be nitrogen and phosphorus deficient. AGS has been successfully cultivated on brewery wastewater [131]. After granulation, high and stable removal efficiencies (i.e., >88%) for organic carbon and nitrogen were achieved at the volumetric exchange ratio of 50% and cycle duration of 6 h. In another study, a 16 h HRT was necessary to provide an adequate starvation phase that benefitted granule stability [88]. Granules were compact with high settling velocities. Nevertheless, further investigations are necessary to focus on decreasing the volumetric exchange ratio or extending the total duration of the SBR cycle in the treatment of high-strength wastewater.
Dairy wastewater

Dairy wastewater is characterized by high concentrations of TSS and particulate substrates such as pieces of cheese, coagulated milk, curd fines, milk film, flavouring agents, and other impurities, such as soil and sand, that enter the sewer system during washing or packaging. High removal efficiencies in terms of COD and nitrogen were obtained at steady-state with mature AGS [132]. However, it was noted that in certain operating conditions, the amount of TSS in the effluent significantly increased. The presence of TSS in the effluent was strongly affected by either the length of the withdrawal period or by the particulate COD applied to biomass ratio. Effluent TSS improved when the withdrawal time was decreased and the pCOD applied to MLVSS ratio was decreased.

One study observed that the maximum OLR was 4 kg-COD/(m$^3$•d) for stable granulation [133]. It was also observed that process efficiency, in terms of VER and cycle duration, was limited by the development of adequate feast and famine conditions [134]. The presence of slowly-biodegradable, organic carbon led to extended feast phases since endogenous respiration rates were not observed during a single SBR cycle. Thus, granules always showed structural deficiencies at high OLR due to filamentous outgrowths on aerobically hydrolyzed organic matter. Decreasing the VER, extending the total duration of the SBR cycle, as well as incorporating an anaerobic hydrolysis period prior the AGS reactor could therefore enhance process efficiency of high TSS wastewaters.
Fish-canning wastewater

Wastewater generated by fish-canning industries are characterized by high salinity (i.e., up to 150 g-NaCl/L), high COD, and TN loads, as well as by a large amount of TSS. Biological treatment of fish-canning wastewater must therefore resist osmotic pressures, generated by high salinity, on cellular membranes that could cause plasmolysis. It has been suggested that shortcut nitrification-denitrification could be the most effective solution for nitrogen removal in saline wastewater. The operational approaches that are generally implemented to inhibit the NOB activity, including high temperature, low DO, decoupled SRT, and high free ammonia concentration are not necessary since salt concentration higher than 20 g-NaCl/L is enough for NOB inhibition [135].

In one instance AGS was used to treat fish-canning wastewater with up to 30 g-NaCl/L [135]. The AGS withheld saline conditions and achieved complete organic matter removal with the presence of nitritation-denitrification activities. The ammonia was mainly oxidized to nitrite and the efficiency of denitrification process was limited to values between 20% and 55%. Another study successfully achieved halotolerant AGS by increasing the salinity in the influent wastewater stepwise [136]. High salt concentrations (i.e., >20 g-NaCl/L) lead to the accumulation of nitrite, most likely due to the inhibition of NOB. The same study reported that simultaneous nitritation-denitrification was successfully sustained at salinities up to 50 g-NaCl/L. The TN concentration in the effluent was less than 10 mg/L at salinities up to 50 g-NaCl/L, revealing a removal efficiency over 95%. However, nitritation efficiency collapsed above 50 g-NaCl/L, likely because oxygen diffusion within the bulk liquid was limited by salt concentration. In contrast, organic matter removal was not affected by salinity, but rather by the OLR.
Both COD and BOD removal efficiencies were over 90%, and specific analyses carried out on the treated wastewater revealed that aerobic granules were able to remove more than 95% of the particulate organic matter. Salinity does, however, significantly influence the physical properties of the aerobic granules [137]. Particularly, with increasing salinity the EPS structure was gradually modified. The fraction of EPS that was not-bound increased with salinity, while the amount of proteinaceous EPS decreased. The modification of the physical structure was more noticeable at salinity above 50 g-NaCl/L, when the granules irreparably lost their stability.

**Palm oil wastewater**

Palm oil production has increased rapidly because of its multiple uses in both food and other industries, especially in the South-East Asia. Direct discharge of palm oil mill effluent into the environment is forbidden due to the high COD concentrations of up to 50,000 mg/L, typically with 50% as BOD. AGS was successfully cultivated at different OLRs ranging between 1.5-3.5 kg-COD/(m³•d) [138]. Different volumetric loading rates led to differences in morphology and structural features of the AGS, in which higher loading rates promoted the formation of larger and looser granules. An interesting observation was that low bacterial speciation was observed in the reactor with high OLR. The bacterial population in the AGS moved away from its initial population representing a permanent change. The authors stated that the shift in microbial population was strictly related to operating conditions. Indeed, it is likely that at high organic loads, the greatest demand for electron acceptors is such that fast-growing, heterotrophic microorganisms have many competitive advantages over slower-growing, autotrophic aerobes.
Winery wastewater

Winery wastewater has been treated in an AGS-SBR [124]. Winery wastewater is characterized by high variability of its organic content. To take into account the fluctuations in organic load, a system to automatically control the HRT based on the anticipated organic load was developed. The relationship between the slope of ORP and DO profiles was used to apply a proper ratio of feast to famine conditions by changing the HRT. At higher loads, a longer HRT was applied to develop an adequate famine phase for the longer feast phase applied. Optimal conditions were achieved when the famine phase was twice as long as the feast phase (i.e., famine phase of 67% the aerated period). Under such operating conditions, the AGS-SBR reached 95% removal of organic matter under a wide range of organic matter concentrations (i.e., 1 to 8 g-COD/L). Furthermore, the AGS-SBR showed high stability and a rapid response to sudden changes in the COD influent concentration.

Petrochemical and oily wastewater

Petrochemical and oily wastewaters are characterized by poor biodegradability due to the presence of hydrocarbons, metal salts, sulphides, phenol, and other substances. Aerobic granules, previously cultivated on synthetic wastewater, were gradually adapted to petrochemical wastewater by gradually increasing the proportion of petrochemical wastewater in the influent [53]. Following the addition of petrochemical wastewater, granule settleability became poor and the EPS matrix began to weaken. When petrochemical wastewater was 100% of the influent, granule settleability and the removal of pollutants deteriorated sharply. The authors observed that adding additional co-metabolites (e.g., propionate) to stimulate the degradation pathways was an effective way to ensure effective removal of nutrients and maintain compact structure of aerobic granules.
It was observed that biomass fed with saline, oily wastewater generated from the washing of oil tankers (i.e., slop) granulated faster compared with conventional synthetic wastewater [54]. However, the granules appeared slightly unstable and more susceptible to breaking. The combined effect of salinity and hydrocarbons caused the inhibition of the autotrophic biomass, with the consequence that nitrification was absent. Hydrocarbons were removed by a physical-biological mechanism, involving an initial adsorption stage followed by the biological degradation of adsorbed hydrocarbons. The adsorption of hydrocarbons within the pores of granules provided longer contact time between microorganisms and pollutant, thereby ensuring the degradation of recalcitrant compounds. Total petroleum hydrocarbons were successfully removed at steady state with removal of over 90%.

The feasibility of treating petrochemical wastewater characterized by high TOC and ammonium nitrogen concentrations (i.e., up to 490 and 630 mg/L, respectively) was evaluated [126]. It was observed that aerobic granules cultivated from non-acclimated biomass irreversibly deteriorated when the percentage of petrochemical wastewater in the influent was raised to 30%. More stable granulation was observed when biomass acclimated to toxic substances was used as inoculum. The same authors stated that due to higher USAV, and to the different sludges used as inoculum, the aerobic granules exhibited a better physical structure, as well as a higher removal efficiency in terms of nitrogen and total organic carbon. In a recent study, two different reactor configurations to treat petrochemical wastewater by means of AGS were evaluated [139]. Particularly, a continuously-aerated SBR with a feast/famine regime and a SBR operated with an anaerobic feast/aerobic famine strategy were examined. The authors observed that cultivation of aerobic granules with petrochemical wastewater was possible, confirming what was previously observed [54]. Furthermore, the feeding regime did not affect the granule formation.
Landfill leachate

Landfill leachate is one of the most significant contamination problems at landfill sites. Leachate contains salts, heavy metals, organic matters, and xenobiotic organic compounds [140]. In addition to xenobiotic organic compounds, landfill leachate is characterized by high ammonium nitrogen loads [141]. An AGS-SBR was studied for landfill leachate treatment [142]. The results demonstrated that COD removal rates decreased as the influent ammonium concentration increased. Concerning nitrogen, it was found that the removal mechanism depended on the influent concentration. When the TAN concentration in the landfill leachate was 366 mg-NH₄-N/L, the dominant nitrogen removal process in the AGS-SBR was SND. Under ammonium concentrations of 788 mg-NH₄-N/L, nitrite accumulation occurred, and the accumulated nitrite was reduced to nitrogen gas by the shortcut denitrification process.

When the influent TAN increased to 1105 mg-NH₄-N/L, accumulation of nitrite and nitrate persisted throughout the cycle, and the removal efficiencies of ammonium decreased to 40%. Similar results were also obtained by another study, which pointed out the decisive role of ammonium concentration in TN removal [143]. The organic and nitrogen removal efficiencies decreased when ammonium concentration increased. For this reason, a specific ammonium pre-treatment must be applied before AGS application. Furthermore, it was emphasized that the start-up phase impacts granulation. It was suggested that the initial cultivation of granules should be carried out with small increments of leachate, in order to optimize the bacteria acclimation and specification. Others have demonstrated that AGS was able to treat a mixture of wastewater with landfill leachate at high ammonium concentration of 900 mg-NH₄-N/L [144]. The COD removal was highly dependent on the ammonium concentration in the influent and on the biodegradability of the organic matter.
Wastewater contaminated by emerging micro-pollutants

Many chemicals used in everyday pharmaceutical and personal care products are among the most commonly detected compounds in surface waters throughout the world. Such compounds may have significant biological effects, such as enzyme inhibitors, modifiers of cell to cell communication, disrupters of membrane transport, and processes for energy generation [145]. Continued exposure to antibiotics can enhance the selection of resistant bacterial strains in the environment [146]. AGS was successfully used for the treatment of aqueous streams containing antibiotics and several pharmaceutical and personal care products. It was demonstrated that conventional AGS was not able to remove 2-fluorophenols [145].

Applying bio-augmentation with a specialized bacterial strain, a strain that was able to degrade 2-fluorophenols, resulted in full degradation of the compound by the AGS. Overall, the AGS-SBR was proven to be robust, exhibiting a high performance after bio-augmentation with the 2-fluorophenols degrading strain. The performances of an AGS reactor treating synthetic wastewater containing fluoroquinolones, a class of antibiotics, was also studied. No evidence of fluoroquinolones biodegradation was observed, but the compound did adsorb to the AGS and was gradually released into the medium in successive cycles after fluoroquinolones were removed from the influent. Overall, neither COD nor ammonium removal were affected during the shock loadings of fluoroquinolones.
The effects of fluoxetine on an AGS reactor were investigated, and it was noted that
fluoxetine adsorption/desorption to granules continuously occurred [147]. Fluoxetine shock loads
did not show a significant effect on the removal of COD or ammonium, whereas a significant
decrease in denitrification removal efficiency was observed. Moreover, a shift in microbial
community occurred probably due to fluoxetine exposure, which induced
adaptation/restructuring of the microbial population. Similarly, changes in the microbial profiles
with time in response to the presence of pharmaceuticals and personal care products were noted
[148]. Initially, pharmaceutical and personal care products addition negatively affected granular
quality, but after microbial adaptation, the system recovered and effectively removed certain
pharmaceutical and personal care products.

AGS in a CFR

General considerations

Operational considerations for an AGS-CFR are essentially the same as considerations
for operation in SBR. There should be enough anaerobic contact of fresh wastewater with
biomass, promoting as much anaerobic uptake of soluble organics as possible. For the same
reason, pre-treatment methods to remove particulate organic matter, should be considered such
that the characteristics of wastewater supplied to the AGS-CFR are able to be converted to
readily bioavailable substrate and then to internal storage products by accumulating
microorganisms.

Granulation of biomass in continuous flow would require representation of what is
known from granulation in SBRs, as well as from the full-scale Nereda® process. It has widely
been suggested that SBRs, which cycle from high to low substrate concentrations over time,
benefit the formation of AGS [31]. Due to the nature of plug-flow CFRs, where a mass of mixed
liquor travels along the length of the reactor, the changing substrate gradients with reactor length would be representative of SBR operation. However, conventional plug-flow BNR processes have numerous recycle lines (i.e., at least one internal recycle and return activated sludge) which dilute substrate concentrations and therefore repress the substrate gradient experienced by the biomass. It would therefore be important to minimize the number of recycle lines as well as the quantity of flow recycled (i.e., no internal recycle, and return activated sludge no more than 100% of the influent flow).

It is well known that the breakthrough of organic substrate in bulk solution to aerobic conditions negatively impacts AGS stability in SBRs. This could be due to many reasons, including the overgrowth of filamentous microorganisms on the surface of granules or an accumulation of particulate matter. Both cases lead to mass transfer limitations and therefore increase the potential for granule disintegration [44], [83]. Thus, it would be important to consider designing the plug-flow CFR configuration such that maximum anaerobic utilization of substrate occurred. This could be done by increasing the volume of the anaerobic zone to account for up to one third of the reactor volume. Under such conditions the hydrolysis of complex, biodegradable influent substrate, as well any residual particulate matter carried over from the return activated sludge line, would be enhanced leading to more anaerobic substrate utilization. Overall, anaerobic substrate utilization promotes uniform growth across the granule and contributes to lower sludge yields.
Allowing for transition from an anaerobic contact phase to an aerobic zone with high oxygen uptake rate, and then a second aerobic zone with low oxygen uptake rate, should promote granulation, if a bulk of the organic carbon is converted to internal storage product in the anaerobic zone. The purpose would be to prevent conditions where fast-growing, aerobic heterotrophs would be allowed to proliferate in bulk solution, which could lead to operational problems. As well, the transition from a high-rate aerobic zone to a low-rate aerobic zone allows for the microbial community to cycle through a phase of EPS production followed by EPS consumption. Therefore, CMR conditions should be avoided in the design of CFRs. Rather, plug-flow or a series of at least two aerobic tanks would be desired.

To move from a hybrid-AGS system (i.e., flocculent biomass and granules) to a completely granular system there must be some degree of selective wasting. Essentially, the SRT of the light biomass fractions (i.e., free-living cells, flocculent biomass) must be decoupled from the SRT of granules and controlled at completely different ends of the spectrum. At 20 °C the minimum SRT for aerobic heterotrophic microorganisms approaches 9 h [86]. Thus, to effectively out-select all free-living, heterotrophic microorganisms in suspension the aerobic SRT of the light biomass fraction would have to be 9 h at 20 °C. In practice, this may be too aggressive an SRT and it is more likely that the range of SRT for the selectively wasted light fraction of biomass would be 1-6 d. On the other hand, previous lab-scale results as well as modelling estimations have demonstrated that the optimum SRT of granules should be controlled at 20 to 50 d. At long granule SRT, desirable metabolic pathways with low yields are selected for, as well as a higher fraction of PAO to OHO [43], [94].
The selective wasting device should be designed with flexibility to allow for diurnal variation in flow, as well as changing flows as population in the sewer catchment increases. For this reason, hydrocyclones are not the most desirable device since the selection pressure in a single hydrocyclone would change with flow. Likewise, physical screening (i.e., 0.2-0.5 mm) is unlikely to provide a robust solution since granules may be subjected to breakage and not necessarily complete disintegration. A much better selective wasting device would decouple the selective wasting criteria from flowrate such that additional operational parameters could be controlled to add system flexibility.

The final consideration for AGS in continuous plug-flow would be the degree of variability allowed. Wastewater varies with season as well as time of day. In fact, in smaller communities the peak hour flow can be three times the average daily flow. With variable flow also comes variable load. Since AGS is sensitive to the breakthrough of organic substrate to the aerobic zone, design should consider minimizing the risk of extended “feast” phases [45].

Objectives

The goal was to develop a continuous flow reactor for aerobic granular sludge that would be suitable for retrofit into existing wastewater treatment facilities where sequencing batch reactors could not be implemented. To attain the goal, several items had to be addressed, including: 1) the suitability of aerobic granular sludge to treat low-strength, municipal wastewater; 2) the importance of hydrodynamic shear force for aerobic granular sludge stability; 3) the role of organic loading rate on aerobic granular sludge stability and performance; 4) the impact of selective wasting on granulation in continuous flow; and 5) the impact of reactor configuration on granulation in continuous flow. Thus, the objectives were to address these items before elucidating design and operational parameters for an AGS-CFR.
Bibliography


[9] D. Rosso *et al.*, “Oxygen transfer and uptake, nutrient removal, and energy footprint of


2013.


2015.


Chapter 2: AGS-SBR treating low-strength wastewater at low hydrodynamic shear force

Introduction

Aerobic granular sludge has been described as one of the most advanced biotechnologies for wastewater treatment [1]. Lab-scale experiments achieved steady-state COD removal of 95%, TIN removal of 94%, and TP removal of 98% [2]. Furthermore, several full-scale AGS-SBR systems treating municipal and industrial wastewaters achieved low effluent nitrogen and phosphorus with up to 60% reduction in energy consumption compared to conventional processes [3], [4]. However, there remained marked scarcity of information regarding treatment of low-strength wastewaters. Especially, drivers behind granule formation, granule stability, and nutrient removal, especially simultaneous nitrification-denitrification SND, when treating low-strength municipal wastewaters were not well described.

Suitability of AGS for low-strength wastewater was an extremely divisive topic in the literature. Some studies suggested that granulation on low-strength wastewater was not possible, while other studies have suggested that increased famine periods, which would suggest lower OLR and therefore low-strength wastewaters, were beneficial for granule stability [5]–[7]. The following summarized some of the findings regarding AGS treating low-strength wastewater. In one case municipal wastewater was pre-fermented with supplemental sucrose addition, thereby increasing the OLR from 1.4 kg-COD/(m³•d) to 2.5 kg-COD/(m³•d), in order to achieve granulation [8]. Without pre-fermentation and sucrose addition granulation was not observed. Another study successfully granulated on low-strength wastewater with 340 mg-COD/L at an OLR of 1.4 kg-COD/(m³•d) [9].
However, that study used a synthetic wastewater which would have been more biodegradable than municipal wastewater. Complete degranulation was observed by others when substrate supplied to an AGS-SBR was switched from an acetate based solution with 1000 mg-COD/L to municipal wastewater with an average COD concentration of 480 mg/L [10]. The same study reported regranulation 30 d after degranulation, thereby suggesting that low-strength wastewater could generate stable AGS after a period of acclimation. Lastly, another study reported successful and stable granulation at various OLR, but indicated that removal efficiency increased as cycle time decreased from 450 min to 200 min (i.e., OLR more than doubled) [11].

This contradiction within the literature regarding AGS treating low-strength wastewater confounded potential applications for AGS processes. Thus, the objective of this study was to examine the development and stability of AGS and measure kinetic rates of nutrient removal while treating low-strength wastewater, without acetate or propionate source, at saturated DO. High DO was used to represent the worst case for SND, as observed by other authors [12]. Biomass characteristics including observed yields, whole sludge SRT, and the diameter of AGS particles were estimated.

Methods

Reactor design and operation

The AGS process was conducted in SBRs with working volume of 4 L at ambient temperature (i.e., 20±1 °C). Three reactors were constructed from 4” (i.e., 10.16 cm) diameter acrylic tube welded to square acrylic sheet base. Access ports were connected to 3/8” (i.e., 0.9525 cm) ID tube at 1 L intervals. A 50% VER was maintained by discharging via the 2 L access port. Thus, the top 2 L of supernatant were discharged in less than 5 min following a 3 min settling period. The HRT was 6 h. Influent was delivered from the bottom through a 3/32”
(i.e., 0.2382 cm) tube over a 1 h, anaerobic period without mixing. This resulted in upflow fluid velocities of 0.41 cm/min relative to the perpendicular cross-sectional area of the reactors. The remaining time was utilized for aerobic reaction, where air was supplied by magnetic piston pumps at a controlled rate of 2 L/min. The air was delivered through three, 3 cm diameter spherical stone diffusers located on the reactor floor. The resulting USAV was 0.41 cm/s.

Cycle duration was 3 h (i.e., 180 min) including 60 min anaerobic feeding, 110 aerobic reaction, 3 min settling, 5 min discharging, and 2 min idling before the next cycle. Sludge was selectively wasted during settling, with sludge not settling within the allotted 3 min having a SRT equal to the 6 h HRT. Cycles were controlled via outlet timers (Traceable Digital Outlet Controller, Fisher Scientific, CA). Dissolved oxygen concentrations in the reactors reached saturation 5 min after aeration began and remained at saturation for the duration of the aerobic period.

The pH was monitored (Eutech pH Controller, Thermo Scientific, USA) and maintained at less than 7.30±0.05 during the aerobic reaction phase using sulfuric acid dosed by peristaltic pumps (Masterflex, Cole-Parmer, CA). Dissolved oxygen concentrations reached saturation within 5 min of aeration and were maintained at saturation during the entire aerobic phase. Peristaltic pumps were also used for the influent and effluent. All reactors were seeded with full-scale biological nutrient removal sludge from the West End Water Pollution Control Centre in Winnipeg, CA, to an initial solids concentration of approximately 3 g-TSS/L. There was no operational difference between the three reactors (i.e., R1, R2, and R3).
A low-strength proteinaceous synthetic wastewater composed of beef and yeast extracts (Sigma Aldrich), ammonium chloride, and potassium phosphate dibasic was supplied to each of the reactors from a common feed tank. Wastewater characteristics were 340±30 mg/L COD (n = 49), 42±5 mg/L TN (n = 48), and 7±1 mg/L TP (n = 48), on average. Wastewater supply lines were disinfected regularly with diluted sodium hypochlorite solution (Javex) and wastewater was stored for 3 d at 4 °C. A heat exchanger in the supply line raised wastewater temperature to 20 °C before introduction to the SBR.

**Analytical methods**

Influent and effluent samples were analyzed for COD, sCOD, TN, and TP (TNTplus, Hach, CA), TAN, nitrite, nitrate, and OP (Lachat QuikChem 8500, HACH, CA). Samples for MLSS, MLVSS, and effluent TSS and VSS were analyzed according to Standard Methods [13]. Individual samples were quantified in triplicate to increase representation of analysis. Kinetic tests examined sCOD, TAN, nitrite and nitrate nitrogen, and OP during regular operational cycles. A SteREO Discovery (Zeiss, DE) was used for microscopic analysis. ImageJ software was used to assess the cross-sectional area and DF of AGS (i.e.,
Appendix A). The cross-sectional area was used to compute $D_s$. All samples that required filtration were run through medium porosity Q5 filter paper (Fisher Scientific, CA). One-way analysis of variance (VassarStats, USA) was used to determine if steady-state data was significantly different (i.e., $\alpha = 0.05$) between R1-R3 (i.e., Appendix B).
The OLR was calculated according to Equation 1. The F/M ratio was determined based on Equation 2. Specific wasting rates were determined according to Equation 3. At steady-state the whole sludge SRT was estimated by taking the inverse of $R_{WAS}$ – Equation 4. Estimates of $Y_{net}$ were determined according to Equation 5. Example calculations for $r_{max}$ are presented in Appendix C.

**Equation 1**

$$OLR = \frac{Q_{influent} \cdot COD_{influent}}{V_{reactor}}$$

**Equation 2**

$$F/M = \frac{Q_{influent} \cdot COD_{influent}}{MLVSS_{reactor} \cdot V_{reactor}}$$

**Equation 3**

$$r_{WAS} = \frac{MLSS_{reactor} \cdot V_{reactor}}{Q_{influent} \cdot TSS_{effluent}}$$

**Equation 4**

$$SRT = \frac{1}{r_{WAS}}$$

**Equation 5**

$$Y_{net} = \frac{Q_{influent} \cdot TSS_{effluent}}{Q_{influent} \cdot COD_{influent}}$$
Results

*Biomass characteristics and quantities*

Biomass in R1-R3 increased from an inoculum concentration of 3 g-VSS/L to their stabilized values. After 40 days of operation the MLVSS in R1-R3 stabilized and ranged from 4 g/L to 7 g/L – Figure 3. The highest MLVSS content was observed in R3 while R1 and R2 had MLVSS ranging from 4-5 g/L. During steady-state the difference in MLVSS content between R1 and R3 (i.e., $P = 0.0033$), as well as R2 and R3 (i.e., $P = 0.0141$), were determined to be significantly different even though operational conditions between R1-R3 were the same. The steady-state MLVSS concentration was 4.2±0.8 g/L in R1, 5±1 g/L in R2, and 7±1 g/L in R3.

*Figure 3 Steady-state MLSS and MLVSS in R1-R3. Error bars represent one standard deviation.*
Effluent VSS in R1 during steady-state was 80±30 mg/L, while it was 60±20 mg/L in R2 and 50±10 mg/L in R3 – Figure 4. It should be noted that higher than usual effluent solids (i.e., usual being <25 mg/L with clarification) observed in R1-R3 were because selective wasting was performed simultaneously with effluent discharge. Full-scale application would require an independent selective wasting stage to avoid effluent solids that would exceed discharge requirements. Typically, <25 mg/L of effluent TSS is specified for municipal wastewater discharge. The difference in steady-state effluent VSS between R1-R3 was determined to be significantly different (i.e., P = 0.0021).

Figure 4 Steady-state TSS and VSS in the effluents of R1-R3. Error bars represent one standard deviation.
The ratio of VSS to TSS in R1-R3 biomass was similar throughout the study. The average VSS to TSS ratio in R1 was observed to be 0.78±0.03, while the VSS to RSS ratio in R2 and R3 were observed at 0.80±0.05 and 0.78±0.04, respectively, during steady-state. The difference in VSS to TSS ratios between R1-R3 during steady-state was not significant (i.e., P = 0.4856). Interestingly, the ratio of VSS to TSS in the effluent of R1-R3 during steady-state was observed to be higher than in the reactors. For instance, the VSS to TSS ratio in R1 effluent was 0.88±0.09, in R2 it was 0.92±0.08, and in R3 it was 0.89±0.09.

Although the difference in effluent VSS to TSS ratios was not significantly different between R1-R3 (i.e., P = 0.5250), there was a significant difference in the VSS to TSS ratio between the biomass in each reactor and the effluent solids from that reactor (i.e., P = 0.0195 for R1; P = 0.0011 for R2; and P = 0.02 for R3). A higher fraction of inert material in the reactor solids (i.e., lower VSS to TSS ratio) was expected in the MLSS since granular sludge was known to accumulate inorganic precipitates [14] and also because inorganic particles settle faster and would not be washed out by selective wasting. Furthermore, the fraction of selectively wasted biomass would be expected to have high VSS to TSS ratios because the low SRT of 6 h applied to slowly settling biomass would result in a more active fraction of slowly settling biomass since less inert solids would accumulate [15].
A mature granular sludge had developed 40 d after inoculation with a suspended biomass culture. The AGS remained stable for the entire 120 d of reactor operation, thereby demonstrating that low-strength wastewater could promote the development and stability of AGS. Previous studies observed degranulation at F/M below 0.4 g-COD/(g-VSS•d) with an influent COD concentration, as yeast extract, of 400 mg/L [5]. Individual reactor F/Ms in this study were 0.33 g g-COD/(g-VSS•d) in R1, 0.29 g g-COD/(g-VSS•d) in R2, and 0.18 g-COD/(g-VSS•d) in R3 during steady-state with an influent COD concentration of 340 mg/L. All three reactors maintained stable granular sludge below 0.4 g-COD/(g-VSS•d), thereby proving that AGS could be operated at lower F/M without degranulation. These results agreed with other studies that have recommended increased periods of famine to benefit AGS stability [6], [7].

There was a distinct increase in granule diameter for R1-R3 during start-up. Average D_F and D_S were quantified for biomass in R1-R3 – Figure 5. During steady-state, average D_F was quantified as 1.3±0.3 mm in R1, 1.3±0.3 mm in R2, and 0.9±0.2 mm in R3. Similarly, average D_S were 1.0±0.2 mm in R1, 1.0±0.3 mm in R2, and 0.7±0.2 mm in R3. There was no significant difference in the average diameters of granules in R1-R3 during steady-state. The ratio of D_F to D_S was determined to be 1.3 in R1-R3. The larger was the ratio of D_F to D_S, the more irregular were the particles. In general, granules appeared uniform with slight surface irregularities. These results suggested that the degree of irregularity in R1-R3 was similar since the ratio of D_F to D_S was determined to be the same.
Biomass growth characteristics

Biomass growth kinetics were observed via \( r_{\text{WAS}} \) and \( Y_{\text{net}} \). All reactors experienced similar trends in \( r_{\text{WAS}} \), with higher values and variability observed during start-up (i.e., the first 30 d from inoculation) and lower, less variable values for the remaining time and during steady-state. Steady-state \( r_{\text{WAS}} \) were determined to be \( 0.08 \pm 0.01 \) g-VSS/(g-VSS•d) in R1, \( 0.05 \pm 0.01 \) g-VSS/(g-VSS•d) in R2, and \( 0.024 \pm 0.004 \) g-VSS/(g-VSS•d) in R3. Like MLSS in R1-R3 and TSS in the effluent from R1-R3, \( r_{\text{WAS}} \) was determined to be significantly different between R1-R3 during steady-state (i.e., \( P = 0.0003 \)). However, the reason for this variation cannot be concluded within the scope of the acquired data.
During steady-state the inverse of $r_{WAS}$ could be assumed as equal to the whole sludge SRT. Although granules were not specifically wasted it was reasonable to assume that at steady-state the decay material from microorganisms within the granule should have been in equilibrium with the decay material in solution (i.e., selectively wasted material would have included decay material from the granules at steady-state). Whole sludge SRT during steady-state was determined to be $14\pm3$ d for R1, while it was determined to be $20\pm4$ d and $42\pm6$ d for R2 and R3 respectively – Figure 6. Steady-state whole sludge SRT, like $r_{WAS}$, were determined to be significantly different between R1-R3 (i.e., $P < 0.0001$) even though there was no difference between reactor operation. Steady-state $Y_{net}$ for R1 was $0.25\pm0.06$ g-VSS/g-COD while $Y_{net}$ for R2 and R3 were $0.20\pm0.06$ g-VSS/g-COD and $0.13\pm0.03$ g-VSS/g-COD, respectively. The difference in steady-state $Y_{net}$ between R1-R3 was determined to be significantly different (i.e., $P = 0.0175$), although the reason for this difference cannot be determined from this dataset. As expected, even if the significant difference between R1-R3 was not significant, $Y_{net}$ decreased as the whole sludge SRT increased.

**Figure 6** Left – steady-state whole sludge SRT in R1-R3; Right – steady-state $Y_{net}$. Errors bars represent one standard deviation.
Phosphorus removal

Orthophosphate release and uptake rates were quantified during regular operational cycles of R1-R3. Over time, from inoculation, similar trends were detected in specific \( r_{\text{max, OP rel.}} \) and \( r_{\text{max, OP up.}} \). Steady-state \( r_{\text{max, OP rel.}} \) and \( r_{\text{max, OP up.}} \) were closely related between reactors. The steady-state specific \( r_{\text{max, OP rel.}} \) in R1 was observed to be 4±2 mg-P/(g-VSS•h), while in R2 and R3 \( r_{\text{max, OP rel.}} \) was observed to be 5±2 mg-P/(g-VSS•h) and 4.2±0.9 mg-P/(g-VSS•h), respectively – Figure 7. There was no significant difference between \( r_{\text{max, OP rel.}} \) in R1-R3 during this period (i.e., \( P = 0.5309 \)). Thus, unlike MLSS, effluent TSS, whole sludge SRT, and \( Y_{\text{net}} \), the \( r_{\text{max, OP rel.}} \) was not impacted by the factor causing significant differences in the other parameters. Other parameters that were not significantly different between R1-R3 included DF and DS, and the ratio of VSS to TSS in the reactors and in the effluent. Similarly, there was no significant difference in \( r_{\text{max, OP up.}} \) (i.e., \( P = 0.6870 \)). Steady-state \( r_{\text{max, OP up.}} \) were determined to be 5±1 mg-P/(g-VSS•h) in R1, 4±1 mg-P/(g-VSS•h) in R2, and 4±1 mg-P/(g-VSS•h) in R3. By comparison, a study using acetate as the sole carbon source obtained \( r_{\text{max, OP rel.}} \) of 14.7 mg-P/(g-VSS•h) and \( r_{\text{max, OP up.}} \) of 11 mg-P/(g-VSS•h) [10]. The use of less readily bioavailable yeast extract in this study, at a COD:TP ratio of 48.6:1, could explain the lower phosphorus release and uptake rates compared with the other study which used acetate at a COD:TP ratio of 68.8:1.
Figure 7 Left – steady-state rate of OP release; Right – steady-state rate of OP uptake. Errors bars represent one standard deviation.

Nitrogen removal

Ammonium utilization rates and nitrite/nitrate production rates were quantified during regular operational cycles of R1-R3. The observed steady-state $r_{\text{max}, \text{NOx prod.}}$ in R1 was 3.0±0.7 mg-N/(g-VSS•h), while it was 1.8±0.2 mg-N/(g-VSS•h) in R2, and 0.9±0.2 mg-N/(g-VSS•h) in R3 – Figure 8. The difference in $r_{\text{max}, \text{NOx prod.}}$ between R1-R3 was determined to be significantly different (i.e., $P = 0.0002$). Thus, nitrite/nitrate production rates were added to the collective group of parameters that were significantly different between R1-R3 despite operational conditions being constant. Other significantly different parameters included whole sludge SRT and $Y_{\text{net}}$.
During steady-state effluent TAN was near or below the detection limit of 0.2 mg/L in R1-R3. Furthermore, TAN in all three reactors during steady-state was near or below the detection limit only 120 min into a single reactor cycle (i.e., aeration from 60-155 min of a single reactor cycle). Thus, the lower $r_{\text{max, NOx prod}}$ observed were not due to less nitrification activity, but instead due to the presence of SND (i.e., less nitrite/nitrate production was observed because it was simultaneously being removed as nitrogen gas). The SND was quantified, relative to the influent TN load, as $25\pm7\%$ in R1 and $33\pm7\%$ in R2, while it was $44\pm7\%$ in R3. The observed SND aligned with the decrease in nitrite/nitrate production rates. For instance, SND in R3 was observed to be 1.8 times higher than R1 and the observed nitrite/nitrate production rate in R1 was observed to be 3.3 times higher than in R3.
Relationship between nutrient removal and biomass characteristics

These results demonstrated that AGS treating low-strength wastewater achieved low effluent TP. During steady-state, effluent OP averaged 0.2±0.4 mg-P L⁻¹. Furthermore, \( r_{\text{max, OP rel.}} \) and \( r_{\text{max, OP up.}} \) between R1-R3 were similar despite significantly different steady-state MLSS concentrations, whole sludge SRT, \( Y_{\text{net}} \), and \( r_{\text{max, NOx prod.}} \). It was interesting to note that despite almost twice the MLSS in R3 compared to R1 that \( r_{\text{max, OP rel.}} \) and \( r_{\text{max, OP up.}} \) remained similar. This suggested that the composition of MLSS, at least in terms of PAO, remained similar between R1-R3 despite significant differences in other parameters. Also suggesting that the composition of biomass between R1-R3 remained similar was the fact that VSS to TSS ratios of biomass in R1-R3 were not significantly different.

However, there was a significant difference in \( r_{\text{max, NOx prod.}} \) even though reactors were operated under the same conditions. It is important to note that despite significant differences in \( r_{\text{max, NOx prod.}} \) and the observed SND that R1-R3 still removed a significant portion of TN. In fact, TN removal was 57±6% in R1, 61±6% in R2, and 67±6% in R3. The differences in nitrification kinetics and SND suggested that the composition of the biomass was significantly different regarding nitrogen removal pathways. In general, nitrifying microorganisms make up a small percentage of the microbial community in biological treatment systems [16]. The low percentage of nitrifying microorganisms may have explained why other compositional parameters (i.e., phosphorus kinetics and VSS to TSS ratios) of the biomass in R1-R3 were not significantly different, although this statement would only be true if the difference in \( r_{\text{max, NOx prod.}} \) was the result of a shift in the nitrifying population. As previously discussed, however, the difference in \( r_{\text{max, NOx prod.}} \) was most likely due to the onset of enhanced SND and not the result of differences in nitrifying populations.
A difference in granule size could have influenced the amount of SND, and therefore \( r_{\text{max}} \), since larger granules may have provided more anoxic volume below the outer aerobic layer for SND [17]. However, there was no significant difference in the steady-state average \( D_F \) or \( D_S \) between R1-R3. Granules in R1 and R2 had average \( D_F \) of 1.3±0.3 mm while the amount of SND, relative to the influent TN load, was 25±7% in R1 and 33±7% in R2 while granules in R3 were the smallest, with an average \( D_F \) of 0.9±0.2 mm, and the amount of SND was the highest at 44±7%. Thus, biomass with smaller average diameter had enhanced SND, which contradicted the general theory that larger granules would have more anoxic volume and thereby more SND. However, these results do agree with more recent studies that have demonstrated limited denitrification rates in granules larger than 0.45 mm [18].

The observed relationships between SND and other significantly different parameters, which included MLVSS concentrations, whole sludge SRT, and \( Y_{\text{net}} \), alluded to some justification behind the observed differences in nitrification kinetics and SND. It should be noted that within the bounds of this study it was impossible to determine causation, and it could only be concluded that the factor impacting SND between R1-R3 also impacted MLVSS, whole sludge SRT, and \( Y_{\text{net}} \). However, it was possible to speak to the relationship between the parameters that were impacted. In general, there was strong correlation between these parameters and SND – Figure 9. For instance, SND positively correlated with MLVSS (i.e., \( R^2 = 0.9173 \)) and whole sludge SRT (i.e., \( R^2 = 0.949 \)), and negatively correlated with observed yield (i.e., \( R^2 = 1 \)).
Figure 9 Steady-state SND as a function of: MLVSS (top left); whole sludge SRT (top right); $Y_{net}$ (bottom left); and $R_{max, OP\ rel.}$ (bottom right). Errors bars represent one standard deviation.

These relationships suggested that enhanced SND was associated with a larger quantity of more slowly growing microorganisms (i.e., higher MLVSS and SRT, and lower $Y_{net}$). If these interrelationships were coupled with the fact that $R_{max, OP\ rel.}$ and $R_{max, OP\ up.}$ were similar between R1-R3, which meant that more mass, overall, of phosphorus was released/uptaken with
increasing MLVSS, it became apparent that SND positively correlated (i.e., $R^2 = 0.9973$) with $R_{\text{max}}$, $OP_{\text{rel.}}$. The apparent significance in difference between R1-R3’s $R_{\text{max}}$, $OP_{\text{rel.}}$ was skewed by the fact that $R_{\text{max}}$, $OP_{\text{rel.}}$ was a combination of insignificantly different (i.e., $r_{\text{max}}$, $OP_{\text{rel.}}$) and significantly different (i.e., MLVSS) parameters. An interesting extension from this relationship, where a higher mass of PAO was associated with enhanced SND, was that DPAO may have contributed to the enhanced SND. Strictly by association, since there are no quantifiable measures of their activity, it could also be suggested that more DGAO were present. These suggestions would also be supported by the relationships between SND and whole sludge SRT and $Y_{\text{net}}$ since anoxic growth of microorganisms is known to occur at lower yields [19].

**Summary**

Overall, these results demonstrated the suitability of AGS to treat low-strength wastewater (i.e., $C/P < 50$) and assessed kinetics rates of nitrogen and phosphorus removal. Effluent OP averaged $0.2 \pm 0.4$ mg-P/L while effluent TIN averaged $17$ mg-N/L. A mature granular sludge formed within 40 d from inoculation with suspended biomass. The AGS remained stable for the 120 d experimentation period, thereby contradicting previous studies that suggested poor granulation and granule stability at low organic loads [5]. Granulation was stable and successful at an average F/M of $0.25 \pm 0.08$ g-COD/(g-VSS•d). Phosphorus kinetics were not significantly different between R1-R3 but there was a significant difference in nitrification kinetics and SND. Interestingly, TN removal was more sensitive than TP removal. Thus, supplemental carbon may be required to achieve lower effluent nitrogen concentrations.
Bibliography


Chapter 3: Impact of OLR on an AGS-SBR at low hydrodynamic shear force

Introduction

The importance of hydrodynamic shear force, typically quantified by USAV, for AGS processes has been popularized in the literature. Upflow superficial air velocity is simply the airflow divided by the cross-sectional area, perpendicular to the airflow, of the reactor. Consensus is that as shear force increases more compact and stable granules are produced. For instance, granulation was not observed in an AGS-SBR treating acetate based substrate when USAV was less than 1.2 cm/s, while the most dense and stable granules were cultivated at USAV of 3.6 cm/s [1]. Similar results were reported in an AGS-SBR treating acetate based substrate when subjected to USAV of 0.8, 1.6, 2.4, and 3.2 cm/s [2]. In that study, granules were observed at 0.8-1.6 cm/s USAV but they were reported to be unstable, while the most stable and dense granules were reported at USAV of 3.2 cm/s. Other studies have reported stable granules at USAV of 2.1-2.8 cm/s in AGS-SBR [3], [4]. However, none of the studies have provided clear justification as to why higher USAV, and therefore higher hydrodynamic shear force, was beneficial for granulation.
Confounding the role of hydrodynamic shear force on AGS formation and stability are studies that have reported stable granules without significant hydrodynamic shear forces. For example, stable granules were observed in an AGS-SBR treating acetate based substrate when USAV was 0.42 cm/s [5]. However, it should be noted that filamentous growth was observed in the same study when carbon was not completely removed during the anaerobic influent distribution phase. Another study varied hydrodynamic shear force in an AGS-SBR by alternating aeration on and off to maintain DO >2 mg/L [6]. This study unintentionally demonstrated that the importance of hydrodynamic shear force may be linked not only to magnitude, but also to contact time. Based on dichotomy in the literature regarding the importance of high hydrodynamic shear force for AGS, a unified theory on the role of hydrodynamic shear force under different process conditions would benefit future AGS applications and studies.

The mechanism through which high hydrodynamic shear force benefits AGS may be by decoupling SRT of the fast-growing microorganisms that tend to accumulate on the surface of granules from the slow-growing microorganisms fixed within the granular matrix. For example, minimum SRT at 20 °C for OHO can be as low as 9 h when considering their net growth rate under non-limiting conditions based on conventional activated sludge models [7]. Thus, there would always be significant potential for free living, fast-growing microorganisms, to accumulate in suspension and become associated with the surface of granules. This issue of fast-growing microorganisms could even manifest itself when subjected to high washouts in low HRT systems, such as AGS-SBR configurations [8], [9].
Therefore, when the potential for fast-growing microorganisms to propagate in the AGS-SBR is high, which would be the case at higher substrate availability and therefore higher OLR, more hydrodynamic shear force may be beneficial. By scouring fast-growing microorganisms from granule surfaces, so that fast-growing microorganisms may be subsequently washed out via selective wasting, higher hydrodynamic shear force may help stabilize AGS. For these reasons, the objective of this study was to examine the impact of increasing wastewater strength, and therefore the OLR, on granular sludge formation and stability at low hydrodynamic shear force. Physical observations, microscopic analysis, effluent characteristics, and biomass kinetics were recorded for each SBR over a period of three to four months. A mechanism for the role of shear force in granulation was proposed and evaluated based on experimental results.

Methods

Reactor design

The experiment used AGS-SBRs with working volumes of 4 L. Five reactors were constructed from 4” (i.e., 10.16 cm) diameter acrylic tube welded to square acrylic sheet base: 1) low-strength 1 (i.e., LSR1); 2) low-strength 2 (i.e., LSR2); 3) low-strength 3 (i.e., LSR3); 4) medium-strength (i.e., MSR); and 5) high-strength (i.e., HSR). A VER of 50% was maintained by discharging via the 2 L access port. The top 2 L of supernatant was discharged in less than 5 min following a 3 min settling period. Upflow fluid velocities were 0.41 cm/min relative to the perpendicular cross-sectional area of the reactors. The remaining time was utilized for aerobic reaction, where air was supplied by magnetic piston pumps at a controlled rate of 2 L/min. Air was delivered through three, 3 cm diameter spherical stone diffusers located on the reactor floor. The resulting USAV was 0.41 cm/s. Aeration provided complete mixing and generated conditions in which DO reached saturation.
Each reactor operated on a 3 h (i.e., 180 min) cycle which consisted of 60 min anaerobic feeding, 110 min aerobic reaction, 3 min settling, 5 min discharging, and 2 min idling. Cycles were controlled via outlet timers (Traceable Digital Outlet Controller, Fisher Scientific, CA). The pH was monitored (Eutech pH Controller, Thermo Scientific, USA) and maintained at less than 7.30±0.05 during the aerobic reaction phase using sulfuric acid dosed by peristaltic pumps (Masterflex, Cole-Parmer, CA). Peristaltic pumps were also used for the influent and effluent. All reactors were seeded with full-scale biological nutrient removal sludge from the West End Water Pollution Control Centre in Winnipeg, CA, to an initial solids concentration of approximately 3 g-TSS/L.

Feed characteristics

Three different synthetic wastewaters simulating low-, medium-, and high-strength wastewaters were supplied to each of the reactors. Three of the reactors (i.e., LSR1-LSR3) received low-strength wastewater composed of beef and yeast extracts (Sigma Aldrich), ammonium chloride, and potassium phosphate dibasic. The low-strength wastewater characteristics were 340±30 mg/L COD (n = 49), 320±30 mg/L sCOD (n = 49), 42±5 mg/L TN (n = 48), and 7±1 mg/L TP (n = 48), on average. The OLR for LSR1-LSR3 was 1.4 kg-COD/(m³•d).

MSR received medium-strength wastewater composed of sodium acetate anhydrous, ammonium chloride, and potassium phosphate dibasic, which was representative of the substrate employed in previous lab-scale AGS studies [3], [4], [10]. In those previous lab-scale studies, granulation was successful on the synthetic wastewater at high USAV. Characteristics of the medium-strength wastewater were 630±80 mg/L COD and sCOD (n = 37), 80±10 mg/L TN (n = 37), and 13±3 mg/L TP (n = 37). The OLR for MSR was 2.5 kg-COD/(m³•d).
HSR received high-strength wastewater composed of fermenter underflow from the Fort Garry Brewing Company in Winnipeg, CA. The high-strength wastewater contained 1300±300 mg/L COD (n = 42), 1200±200 mg/L sCOD (n = 42), 50±20 mg/L TN (n = 40), and 6±2 mg/L TP (n = 40), on average. The OLR for HSR was 5.2 kg-COD/(m³•d).

Every synthetic wastewater was supplemented with alkalinity as sodium bicarbonate and trace elements [11]. Particulate matter was minimized in the synthetic wastewaters since it has been demonstrated to affect AGS surface properties and would therefore confound results. Specifically, it was observed that particulate matter sorbed to granule surfaces and slowly hydrolysed during aeration, encouraging irregular bumps to form on the granule surface where particulates had been enveloped [12].

Analytical methods

Influent and effluent samples were analyzed for COD, sCOD, TN, TP (TNTplus, Hach, CA), TAN, nitrite nitrogen, nitrate nitrogen, and OP (Lachat QuikChem 8500, HACH, CA). Samples for MLSS, MLVSS, and effluent TSS and VSS were analyzed according to Standard Methods [13]. A SteREO Discovery (Zeiss, DE) was used for microscopic analysis. All samples that required filtration were run through medium porosity Q5 filter paper (Fisher Scientific, CA). Volatile fatty acids were determined by a Breeze 2 HPLC system (Waters, Milford, MA, USA) equipped with an Aminex HPX-87H column using 5 mM H₂SO₄ as the mobile phase, calibrated using a VFA standard 10 mM mixture (Supelco, Bellefonte, USA). Anaerobic sCOD utilization was estimated according to Equation 6. Values for $r$ were estimated according to Equation 7, where $S$ could be COD, TN, or TP.
Equation 6

\[
\text{Anaerobic sCOD utilization} = \left(1 - \frac{s\text{COD}_{\text{end of anaerobic influent distribution}} \cdot V_{\text{reactor}}}{s\text{COD}_{\text{effluent}} \cdot V_{\text{effluent}} + s\text{COD}_{\text{influent}} \cdot V_{\text{influent}} \cdot \text{cycle}} \right) \cdot 100
\]

Equation 7

\[
r = \frac{Q_{\text{influent}} \cdot (S_{\text{influent}} - S_{\text{effluent}})}{\text{MLVSS}_{\text{reactor}} \cdot V_{\text{reactor}}}
\]

Results

General performance

Reactors LSR1-LSR3 maintained effluent sCOD concentrations below 50 mg/L throughout the study. Soluble TN in the effluent consistently decreased during operation, as did mixed liquor suspended solids increase. In LSR3, effluent sTN reached concentrations below 12 mg/L. Effluent sTP stabilized in 20 d for all three SBR receiving low-strength wastewater. There were fluctuations above the average effluent sTP, especially for LSR2. Changes in effluent sTP, as well as sTN and VSS, for LSR2 were attributed to two unique operational failures: 1) aeration failed for at least 48 h on day 61; and 2) feed had stopped for 24 h on day 104. The first failure resulted in decreased nitrification, and consequently decreased effluent nitrate and increased effluent sTN. As nitrification recovered effluent sTN in LSR2 began to decrease and effluent nitrate began to increase. By day 76 the transition back to full nitrification had upset phosphorus removal, likely due to the return of nitrate in the effluent. Cultivation of PAO, as demonstrated by low effluent sTP, correlated with the maturation of aerobic granules in all SBR receiving low-strength wastewater. Effluent VSS stabilized after 40 d and VSS below 30 mg/L in the effluent were only achieved in LSR3. High effluent TSS and VSS in all reactors can be attributed to selective wasting occurring simultaneously with effluent discharge.
MS also achieved consistent effluent sCOD below 50 mg/L for the entire study. Effluent sTN was observed to stabilize early on, with fluctuations in concentration matching the trend in influent TN. Soluble TN in the effluent did not go below 40 mg/L, suggesting poor nitrogen removal capabilities. Phosphorus removal had initially stabilized after 20 d, although effluent sTP was observed to increase after 36 d. Effluent sTP decreased again after 66 d. The trend in effluent sTP matched what was observed in effluent VSS. Effluent VSS peaked above 600 mg/L after 40 d before dropping to 150 mg/L after 60 d. The initial cultivation of PAO matched the appearance of granules in MSR, while the appearance of filamentous growth on the surface of granules aligned with the increase in effluent sTP and VSS.

Effluent sCOD in HSR did not stabilize until day 30. Following stabilization, effluent sCOD remained below 200 mg/L. Both sTN and sTP in the effluent of HSR correlated with TN and TP in the feed. Total nitrogen and TP in the high-strength wastewater were low relative to COD in the influent. Thus, removal of TN and TP in HSR were mainly attributed to assimilation into new biomass. Effluent VSS fluctuated often and stabilized around 300 mg/L. Effluent VSS also peaked after 70 d, reaching 900 mg/L. High effluent VSS in HSR suggested a highly unstable biomass consisting of fast-growing microorganisms.
Biomass quantities and growth parameters

Solids accumulated in the AGS-SBRs over the entire study since the only method of biomass removal was via effluent VSS – Figure 10. Biomass in HSR plateaued at 1.7±0.6 g-VSS/L. The VSS/TSS for HSR was 0.92. Within 10 d the biomass in MSR had stabilized at approximately 2.3±0.5 g-VSS/L. There was a slight increasing trend for biomass in MSR after 50 d, and the VSS/TSS for the entire study was 0.81. Biomass in LSR1-LSR3 did not plateau until 90 d, after which there may still have been an increasing trend in VSS. After 90 d the average biomass concentrations in LSR1-LSR3 were 4.7±0.4 g-VSS/L, 5±1 g-VSS/L, and 8.0±0.3 g-VSS/L, respectively. The VSS/TSS after 90 d was 0.76 in all three AGS-SBR receiving low-strength wastewater.

Figure 10 MLVSS for all AGS-SBRs over the entire study.
In long SRT systems, such as AGS-SBR where granular sludge wasting does not occur, biomass would be expected to accumulate as OLR increased. These results point to the opposite for AGS cultivated in SBR and subjected to low USAV. This phenomenon was caused by the selection pressure applied to select for dense, rapidly settling biomass typical of AGS processes. Given the stochastic nature of solids separation during settling, compounded with USAV not being high enough to sufficiently scour fast-growing microorganisms from granule surfaces, fast-growing microorganisms were able to form a stable population in HSR and MSR despite low HRT of 6 h and therefore 6 h SRT\(_{SW}\). The shift in kinetic nature of the biomass was best demonstrated by \(r_{WAS}\). Values for \(r_{WAS}\) were 1.04 g-VSS/(g-VSS•d) and 0.28 g-VSS/(g-VSS•d) during the last month of operation for HSR and MSR, respectively. Values for \(r_{WAS}\) in LSR1-LSR3 was 0.06 g-VSS/(g-VSS•d), 0.10 g-VSS/(g-VSS•d), and 0.02 g-VSS/(g-VSS•d), respectively, for the last month of operation. At steady-state, a relationship between \(r_{WAS}\) and SRT could be assumed. Thus, the whole sludge SRT during the final month of operation was 0.96 d for HSR, 3.6 d for MSR, 17 d for LSR1, 10 d for LSR2, and 50 d for LSR3.

*Physical appearance and microscopic analysis*

Significant changes in biomass morphology were observed within as little as one week. High solids washout was likely the most influential parameter governing initial changes in the biomass community. Since wastewater characteristics were constant the initial solids washout resulted in F/M peaked at 14 kg-COD/(kg-VSS•d) in HSR and 2.4 kg-COD/(kg-VSS•d) in MSR. Food to microorganism ratios in LSR1-LSR3 peaked 1.4 kg-COD/(kg-VSS•d), 3.9 kg-COD/(kg-VSS•d), and 1.2 kg-COD/(kg-VSS•d), respectively. The F/M was higher in LSR2 because it experienced more solids washout initially. For instance, MLVSS was 0.9±0.2 g/L, 0.3±0.1 g/L, and 1.14±0.03 g/L in LSR1-LSR3, respectively, four days after inoculation.
Previous studies have suggested that high F/M benefits the conversion of flocculent biomass to granules [10]. High F/M during granulation may have allowed for the accumulation of EPS since there would have been less EPS consumption during the aerobic period [14]. Extracellular polymeric substances are known to be an important component of AGS structure. Accumulation of biomass in the AGS-SBR caused F/M to decrease with time, potentially resulting in lower net production of EPS due to aerobic consumption. Lowest observed F/M in HSR and MSR was 1.8 kg-COD/(kg-VSS•d) and 0.76 kg-COD/(kg-VSS•d), respectively. In LSR1-LSR3 the lowest F/M observed were 0.23 kg-COD/(kg-VSS•d), 0.23 kg-COD/(kg-VSS•d), and 0.15 kg-COD/(kg-VSS•d), respectively.

Influent characteristics may have also played a significant role in the initial stages of HSR. Growth was most likely limited by nitrogen and phosphorus concentrations even though there was an abundance of organic carbon available for OHO to grow. Chemical oxygen demand to nitrogen ratio and COD/TP for the high-strength wastewater averaged 50±20 and 300±80, respectively, for the first 40 d. A ratio of 100 COD/10 TN/1 TP was reported to be ideal for AGS treating brewery wastewater, suggesting that the COD/TN and COD/TP were much higher in this study and therefore nitrogen and phosphorus may have been limiting [15]. Mature granulation occurred within 40 d in all but HSR. The physical appearance of biomass, besides colour, was almost indistinguishable between MSR and LSR1-LSR3. Granules in MSR appeared whitish-brown while the low-strength wastewater generated yellowish-brown granules. Large granules were observed in HSR after 2 d. However, after a month HSR was characterized by a viscous, white substance with filamentous microorganisms. The material in HSR was most likely a mixture of EPS and some active cells.
The characteristics of biomass in HSR did not improve for the remainder of the study. Granules with diameters of 0.5-5 mm were observed in all other AGS-SBR, although granule surface properties varied significantly between reactors receiving medium- and low-strength wastewater. Granules in MSR maintained a filamentous surface layer, which was observed one month after inoculation. On the other hand, granules in LSR1-LSR3 appeared smooth and highly compact. On low-strength wastewater LSR3 had the best granules in terms of appearance, while LSR2 had the worst. It should be recalled, however, that there were two distinct operational failures for LSR2 that interfered with stabilization of granular sludge. Therefore, conclusions cannot be directly drawn from the results of LSR2 versus LSR1 and LSR3.

Overall, physical and microscopic analysis of biomass in the five AGS-SBR confirmed the suggested theory for the role of shear in granular sludge formation and stability. Smooth, uniform granules were observed in all three AGS-SBR treating low-strength wastewater even though the applied USAV was low. Granule appearance progressively degraded from LSR to MSR and then to HSR, respectively. Surface fouling was observed in MSR, suggesting that the applied hydrodynamic shear force was not enough to remove rapidly growing, filamentous microorganisms from granule surfaces. These results demonstrated that high hydrodynamic shear forces are not always a necessity for granulation, although they may benefit granular formation and stability depending on wastewater characteristics, such as carbon content.
Specific removal rates and nutrient profiles

Removal of COD, TN, and TP were recorded to compare biomass activity at different OLR. Based on effluent sCOD the HSR removed 62% of COD within the first month. Effluent VSS increased consistently with influent COD load. Lowest effluent VSS and highest MLVSS was observed in LSR3 while the worst effluent VSS and lowest MLVSS was recorded in HSR. Organic carbon removal in HSR stabilized by the second month. Percent COD removal was 89%, 85%, and 85% in the second, third, and fourth month, respectively, based on effluent sCOD. Removal of organic carbon in MSR stabilized quickly. Chemical oxygen demand removal of 92%, 94%, and 94% was observed in the first, second, and third month, respectively, based on effluent sCOD. Organic carbon removal also stabilized quickly in LSR1-LSR3. Percent removal based on effluent sCOD ranged from 87% to 92% in all three low-strength AGS-SBR during the four months of analysis.

Nitrogen and phosphorus removal in HSR were likely carried out entirely by adsorption/assimilation. Biological phosphorous removal was observed in MSR but nitrification was not. The mechanism of nitrogen removal in MSR was therefore entirely based on assimilation, since particulate nitrogen was not present in the feed. Total phosphorous removal in MSR increased consistently from 31% to 69% to 85% in the first, second, and third month, respectively, while nitrogen removal beyond 35% was not observed.
Biological phosphorous removal, nitrification, and denitrification were observed in all three AGS-SBR treating low-strength wastewater. Phosphorous removal during the first month was 71%, 53%, and 87% in LSR1, LSR2, and LSR3, respectively. For the remaining three months phosphorous removal ranged from 85% to 99% in all three AGS-SBR. This range excluded the last month of data for LSR2 because of the operational failures discussed previously. Nitrogen removals of 52%, 42%, and 52% were observed during the first month in LSR1, LSR2, and LSR3, respectively. The range of nitrogen removal for all three AGS-SBR treating low-strength wastewater during the last three months was 50% to 68%. Denitrification was most significant in LSR3, with denitrification having carried out approximately 35% of TN removal during the anaerobic feeding period.

Performance measures for the five AGS-SBR during granulation agreed with results from microscopic analysis, and thus further strengthened the theory for the role of shear in granule sludge formation and stability. Performance in HSR was poor, with undesirable biomass characteristics and high effluent solids contributing to effluent COD. MSR performed better than HSR, although the lack of nitrification in MSR indicated deficiencies in treatment capacity. The layer of filamentous microorganisms on the surface of granules in MSR could have contributed to mass transfer limitations and therefore preferential oxidation of carbon instead of ammonium in the thin aerobic layer of the biofilm. The best performing SBR were LSR1-LSR3, with significant organic carbon removal, nitrification, biological phosphorous removal, as well as some denitrification. Performance measures further demonstrated that granulation does not always require high hydrodynamic shear force, and rather that hydrodynamic shear force may play a more important role in granule formation and stability in the presence of higher OLR to the aerobic zone.
Figure 11 Specific COD, TN, and TP removal rates for each AGS-SBR during the last month of operation (i.e., steady-state). Error bars represent one standard deviation.
Specific COD, TN, and TP removal rates in HSR were highly variable – Figure 11. This can be attributed to fluctuation in effluent VSS and therefore constantly fluctuating MLSS and MLVSS quantities. Specific removal rates in all other AGS-SBR, besides HSR, were much more stable. Removal rates for COD and TP were higher in MSR than in the three AGS-SBR receiving low-strength wastewater. Specific TN removal rates were comparable between MSR and the three AGS-SBR receiving low-strength wastewater. In terms of physical appearance, LSR3 had the most smooth and uniform granules. In terms of performance LSR3 also had the lowest $r_{\text{COD}}$ of 0.16 kg-COD/(kg-VSS•d), $r_{\text{TN}}$ of 17 g-TN/(kg-VSS•d), and $r_{\text{TP}}$ of 3.1 g-TP/(kg-VSS•d). Thus, it can be concluded that slower growing microorganisms with lower specific removal rates contributed to more stable granulation.

Soluble COD in HSR increased during the anaerobic influent distribution phase by up to 25% from what would have been expected if the new feed and previous effluent were combined without considering microbial activity (e.g., biodegradation or uptake of COD) – Figure 12. The most likely explanation was that particulate matter, in the feed or leftover from the previous cycle, hydrolyzed during the anaerobic influent distribution phase. Complete breakdown of particulate organics in HSR during the aerobic phase was unlikely since F/M never dropped below 1.8 kg-COD/(kg-VSS•d). Thus, there could have been leftover particulate COD from the previous cycle or organic products of microbial origin that were hydrolyzed during anaerobic influent distribution. Furthermore, since there was no nitrification in HSR, COD would not have been utilized for denitrification. Therefore, the anaerobic influent distribution phase of HSR operated as a contact/fermentative period.
There was no decrease in sCOD was observed in MSR during the anaerobic influent distribution phase. Organic carbon in the medium-strength wastewater consisted purely of acetate while sCOD remaining at the end of the anaerobic feeding phase was determined to contain approximately 95% acetate by HPLC analysis. Thus, approximately 16 mg-COD/L of acetate was utilized during the anaerobic feeding phase. Phosphorous release in MSR was observed up to 30 min into the aerobic period, with approximately 75% of the mass of phosphorous released during anaerobic feeding – Figure 13. Therefore, although there was PAO activity, the anaerobic influent distribution phase in MSR acted as a poor buffer for sCOD breakthrough to the aerobic zone. The amount of sCOD available to fast-growing microorganisms under aerobic conditions in MSR was enough to sustain an active population. In the case of MSR, it was possible that higher USAV would have improved performance and AGS stability by removing fast-growing microorganisms from the granule surface.
Significant anaerobic sequestration of sCOD from bulk solution was observed in LSR1-LSR3. Specifically, 55% to 70% of the incoming sCOD load was removed from bulk solution during the anaerobic influent distribution. Some sCOD would have been utilized for denitrification of residual nitrates and phosphorous release by PAO, while the remaining sCOD could have diffused from bulk solution into the pore space of granules. Regardless of the mechanism behind anaerobic carbon utilization, the load of carbon utilized in LSR1-LSR3 was enough to prevent overgrowth of fast-growing microorganisms during aeration and therefore maintain stable granules under low USAV. Therefore, high hydrodynamic shear force was not required for stable granulation on the low-strength wastewater, while higher hydrodynamic shear force would most likely have improved the quality of biomass in HSR and MSR.
Summary

It was demonstrated that high hydrodynamic shear force, characterized by high USAV, was not required for granulation. Mature AGS developed within 40 d and was stable for the entire 120 d experiment when an OLR of 1.4 kg-COD/(m³•d) was applied. At OLR of 2.5 kg-COD/(m³•d) granulation was successful, although the granules were characterized by a filamentous surface layer and no nitrification. Furthermore, at OLR of 5.2 kg-COD/(m³•d) granules quickly disintegrated 2 d from inoculation and the biomass took on a white, viscous appearance with filamentous microorganisms. Thus, higher hydrodynamic shear force would likely benefit AGS processes with OLR of 2.5 kg-COD/(m³•d) and higher, since the shear force would act to remove fast-growing, filamentous microorganisms from AGS surfaces. Lastly, it was demonstrated that the most stable AGS had the lowest specific removal rates of COD, TN, and TP. Therefore, it was important to consider that although AGS allows for process intensification, the intensification is predicted by retention of a large quantity of slow-growing microorganisms.
Bibliography


Chapter 4: Controlled SRT stabilizes AGS-SBR under high OLR and low hydrodynamic shear force

Introduction

AGS has provided treatment of wastewaters with up to 60% less energy than conventional technologies [1]–[3]. The benefits of AGS processes result from self-immobilized, spherical-shaped biofilms that allowed for process intensification as well as changes to the metabolic capabilities of biomass [4]. As a result, carbon, nitrogen, and phosphorus can be removed in a single tank with smaller footprint than conventional technologies [5]. Process intensification provided by AGS means that higher OLR than conventional technologies can be utilized. However, there was discrepancy in the literature regarding stability of AGS under high OLR [6]. Furthermore, there appeared to be a lack of methods, beside high hydrodynamic shear force, to provide for AGS stability under increasing OLR.

High OLR would not be an issue for AGS treating municipal wastewaters where the lower range of OLR is actually the problem [7]–[9]. However, there are many industries that produce high-strength, organic wastewaters [10], [11]. Pork (i.e., COD >4000 mg/L), poultry (i.e., COD >1800 mg/L), and red meat (i.e., COD >5900 mg/L) processing plants all produce high-strength, organic wastewaters [10]. Breweries produce many streams of high-strength wastewater, including malting (i.e., COD >500 mg/L), saccharification/fermentation (i.e., COD >6500 mg/L), and packaging (i.e., COD >1500 mg/L) [12]. Similarly, agricultural industries such as piggeries (i.e., COD >950 mg/L) and sugar mills (i.e., COD >3500 mg/L) produce high-strength wastewater [10], [12]. These examples clearly demonstrated potential applications for
AGS at high OLR. Based on the literature and previous lab-scale studies (i.e., Chapter 3), these applications would require a method to maintain AGS stability.

Hydrodynamic shear force may be defined as the force generated due to the friction between the bulk liquid and granules surface. The hydrodynamic shear force has typically been quantified by the USAV, which is calculated as the airflow divided by the cross sectional area of the aerated zone [13]. At high OLR most of the literature agreed that high hydrodynamic shear force was required, although there was discrepancy as to the exact value [14]. For instance, granulation was not observed when USAV was less than 2.4 cm/s at an OLR above 9 kg-COD/(m²•d) while USAV of 3.2 cm/s allowed for OLR of 15 kg-COD/(m²•d) [15]. Similarly, with OLR of 6 kg-COD/(m²•d) unstable granules developed when USAV was less than 2.4 cm/s [16]. With low USAV of 0.41 cm/s AGS was observed to be stable at 1.4 kg-COD/(m²•d), although at 2.5 kg-COD/(m²•d) AGS developed a filamentous surface layer after one month and at 5 kg-COD/(m²•d) no AGS was observed [13]. All these examples demonstrated that although AGS can be stable at low hydrodynamic shear forces, high shear was required to stabilize AGS at high OLR.

The relationship between hydrodynamic shear force and OLR suggested that shear force was required to reduce the quantity of fast-growing microorganisms that cultivated granule surfaces at high substrate availability (i.e., high OLR) [13], [17]. Thus, if the mechanism of hydrodynamic shear force was to selectively remove microorganisms, another mechanism of encouraging granule stability would be to control the granular sludge solids retention time SRT. The objective of this study was to examine the development and stability of AGS with controlled SRT under different OLR, as well as to observe the impact of eliminating controlled wasting of biomass on AGS morphology and performance. All reactors operated under low hydrodynamic
shear force. Reactor performance was quantified based on effluent parameters. Biomass characteristics including net observed yields, nitrogen and phosphorus content of biomass, and SRT were estimated.

**Methods**

*Reactor design and operation*

Experiments were carried out in 4 L sequencing batch reactors (i.e., R1-R3). Air was supplied at 2.5 L/min resulting in upflow superficial air velocity of 0.51 cm/s. Cycle length was 6 h, consisting of 35 min anaerobic influent distribution, 310-312 min aeration, and 4.5 min, 3 min, and 1.5 min of settling in R1, R2, and R3 respectively, before 2 min of effluent discharge. The settling time was different in each reactor to maintain constant critical settling velocity since VER differed between R1-R3. For instance, R1 had 75% VER, R2 had 50% VER, and R3 had 25% VER. Therefore, 3 L, 2 L and 1 L of effluent was withdrawn from R1, R2, and R3, respectively, at the end of each cycle. The critical settling velocity in R1-R3 was 8 cm/min. As a result of different VER, the hydraulic retention time of R1 was 8 h, while it was 12 h for R2 and 24 h for R3. All reactors were seeded with full-scale biological nutrient removal sludge from the West End Water Pollution Control Centre in Winnipeg, CA, to an initial solids concentration of approximately 3 g-TSS/L.

The reactors were supplied with brewery wastewater collected from a local brewery. The raw wastewater was pre-treated in an anaerobic moving bed biofilm reactor [11], and its effluent was used as influent for the AGS reactors. Characteristics of the anaerobically pre-treated brewery wastewater were 1500±200 mg-COD/L, 1300±200 mg-sCOD/L, 140±9 mg-TN/L, and 19.3±0.4 mg-TP/L. Thus, the OLRs for R1, R2, and R3, were 4.5±0.6 kg-COD/(m³•d), 3.0±0.4 kg-COD/(m³•d), and 1.5±0.2 kg-COD/(m³•d), respectively. 450 mL of mixed liquor was
removed from the reactors at the end of the aerobic phase 5 d a week. Therefore, the effective flow of waste mixed liquor was approximately 320 mL/d. The reactors reached steady-state, in terms of MLSS, COD removal, and granule size, within 30 d (i.e., approximately 3 times the SRT) from inoculation and the steady-state period of analysis lasted 30 d.

**Analytical methods**

Influent and effluent samples were analyzed for COD, sCOD, TN, and TP (TNTplus, Hach, CA), TAN, nitrite nitrogen, nitrate nitrogen, and OP (Lachat QuikChem 8500, HACH, CA). Samples for MLSS, MLVSS, and effluent TSS and VSS were analyzed according to Standard Methods [18]. Kinetic tests examined sCOD, TAN, nitrite and nitrate nitrogen, and OP during normal operational cycles. A SteREO Discovery (Zeiss, DE) was used for microscopic analysis. All samples that required filtration were run through medium porosity Q5 filter paper (Fisher Scientific, CA). One-way analysis of variance (VassarStats, USA) was used to determine if steady-state data was significantly different (i.e., $\alpha = 0.05$) between R1-R3.

Organic loading rate was estimated based on Equation 1. The SRT was estimated according to Equation 8. It should be noted that the SRT calculation was for the whole sludge (i.e., wasted mixed liquor and effluent solids). The SRT of particles settling faster than the critical settling velocity of 8 cm/min (i.e., $\text{SRT}_{\text{AGS}}$) could be estimated purely on mixed liquor wasting, which simplified to Equation 9. However, because there would have been some exchange of decayed material from AGS into bulk solution, the true SRT of AGS likely existed between the estimated SRT and $\text{SRT}_{\text{AGS}}$. The $Y_{\text{net}}$ was estimated according to Equation 10. The $N_{\text{MLVSS}}$ and $P_{\text{MLVSS}}$ were estimated by Equation 11 and Equation 12, respectively. The RR was used to assess reactor performance after mixed liquor wasting was stopped and it was determined according to Equation 13.
Equation 8
\[
SRT = \frac{MLSS_{\text{reactor}} \cdot V_{\text{reactor}}}{Q_{\text{WAS}} \cdot MLSS_{\text{reactor}} + Q_{\text{influent}} \cdot TSS_{\text{effluent}}}
\]

Equation 9
\[
SRT_{\text{AGS}} = \frac{V_{\text{reactor}}}{Q_{\text{WAS}}}
\]

Equation 10
\[
Y_{\text{net}} = \frac{Q_{\text{WAS}} \cdot MLSS_{\text{reactor}} + Q_{\text{influent}} \cdot TSS_{\text{effluent}}}{Q_{\text{influent}} \cdot COD_{\text{influent}}}
\]

Equation 11
\[
N_{\text{MLVSS}} = \frac{Q_{\text{influent}} \cdot (TN_{\text{influent}} - TN_{\text{effluent}} - N_{\text{denitrification}})}{Q_{\text{WAS}} \cdot MLVSS_{\text{reactor}} + Q_{\text{influent}} \cdot VSS_{\text{effluent}}}
\]

Equation 12
\[
P_{\text{MLVSS}} = \frac{Q_{\text{influent}} \cdot (TP_{\text{influent}} - OP_{\text{effluent}})}{Q_{\text{WAS}} \cdot MLVSS_{\text{reactor}} + Q_{\text{influent}} \cdot VSS_{\text{effluent}}}
\]

Equation 13
\[
RR = \frac{\text{Removal}_{\text{without SRT}}}{\text{Removal}_{\text{steady-state with SRT}}}
\]
Results

Reactor performance with SRT

Rector performance was based on effluent COD, sCOD, TN and TP – Figure 14. Effluent COD was 280±60 mg/L in R1, 260±60 mg/L in R2, and 220±50 mg/L in R3. The difference in effluent COD between R1-R3 was determined to be significantly different (i.e., P = 0.0400), therefore OLR did have a significant impact on COD removal between R1-R3 (i.e., OLR was 4.5±0.6 kg-COD/(m³•d) for R1, 3.0±0.4 kg-COD/(m³•d) for R2, and 1.5±0.2 kg-COD/(m³•d) for R3). The COD removals, relative to the influent COD of 1500±200 mg/L, were 82%, 83%, and 86% for R1-R3, respectively. Thus, at higher OLR the reactors removed slightly less COD.

Figure 14 Effluent COD, sCOD, TN, and TP in R1-R3. Error bars represent one standard deviation.
Similarly, there was a significant difference in effluent sCOD between R1-R3 (i.e., P <0.0001). Effluent sCOD was 230±40 mg/L in R1, 220±50 mg/L in R2, and 60±40 mg/L in R3. Relative to influent sCOD of 1300±200 mg/L the sCOD removals were 78% in R1, 83% in R2, and 95% in R3. Therefore, OLR also had an impact on the sCOD removal. There was no significant difference in effluent TN (i.e., P = 0.9236) or effluent TP (i.e., P = 0.2874). Thus, the OLR had no impact on TN or TP removals. Effluent TN was 100±10 mg/L in R1, 94±6 mg/L in R2, and 90±10 mg/L in R3. Effluent TP was 14±1 mg/L in R1, 13±2 mg/L in R2, and 13±2 mg/L in R3. Total nitrogen and TP removals were observed to range from 31-33% and 30-35%, respectively, for R1-R3. Effluent TSS was observed to be 90±30 mg/L in R1, 100±100 mg/L in R2, and 130±50 mg/L in R3. The difference in effluent TSS was not significantly different between reactors (i.e., P = 0.38640), and therefore the OLR did not impact effluent TSS.

Trends in sCOD and nitrite/nitrate were monitored over regular operational cycles during steady-state – Figure 15. It should be noted that the sCOD and nitrite/nitrate concentrations at time zero were based on the load of substrate in the effluent from the previous cycle and the load of substrate that would be supplied during the anaerobic influent distribution period. Soluble COD concentrations at time zero were different for R1-R3 because of different VERs (i.e., R1 had 75% VER, R2 had 50% VER, and R3 had 25% VER). Therefore, R1 received more brewery wastewater during anaerobic influent distribution resulting in higher sCOD concentrations than R2 and R3. At time zero the sCOD concentrations were 900±200 mg/L in R1, 700±100 mg/L in R2, and 430±90 mg/L in R3.
Anaerobic influent distribution occurred from time 0-35 min. The 35 min sample had 600±200 mg/L sCOD in R1, 500±90 mg/L sCOD in R2, and 300±60 mg/L sCOD in R3. Relative to the sCOD concentration at time zero, R1-R3 were able to utilize 23%, 23%, and 30% of sCOD, respectively, under anaerobic conditions. The slope of sCOD residual versus operational time changed when sCOD concentrations in the reactors was approximately 250 mg/L. Thus, it could be suggested that the more bioavailable fraction of sCOD was removed by the time sCOD in the reactor reached 250 mg/L. In R1 the sCOD reached 250 mg/L after 75 min (i.e., 40 min into aeration), while R2 reached 250 mg/L sCOD after 60 min (i.e., 25 min into aeration), and R3 reached 250 mg/L sCOD after 45 min (i.e., 10 min into aeration).
The time which bioavailable substrate is present in bulk solution during aeration has been referred to as the feast period [19]. The duration of the feast period in R1-R3 was estimated to be 40 min, 25 min, and 10 min, respectively. Thus, the ratio of feast conditions (i.e., time during aeration with bioavailable substrate in bulk solution) to famine conditions (i.e., time during aeration without bioavailable substrate in bulk solution) were 0.13 in R1, 0.08 in R2, and 0.03 in R3. These ratios would be considered sufficient for generation of stable AGS [20]. The amount of sCOD removed by R1-R3 during the feast period was 400±100 mg/L, 300±100 mg/L, and 100±100 mg/L, respectively. Based on the different feast to famine ratios and quantity of sCOD available during the feast period, it could be concluded that the OLR had an impact on the feast conditions in R1-R3, with regards to both the time of feast conditions and the quantity of substrate available during feast conditions.

During steady-state operation, there was high variability in effluent nitrite and nitrate. The variability was quantifiable through high standard deviations for nitrite and nitrate concentrations depicted in Figure 16. Regardless, the presence of some nitrite and nitrate suggested that denitrification may have been occurring during anaerobic influent distribution and therefore prevented full anaerobic conditions from developing. Based on the nitrite/nitrate concentrations at the beginning and end of the anaerobic influent distribution, it was determined that R1 denitrified 11±14 mg-N/L, R2 denitrified 17±74 mg-N/L, and R3 denitrified 7±45 mg-N/L. Assuming a ratio of 6.6 g-COD/g-N required for denitrification of nitrate [21], which was conservative since not all the nitrite/nitrate was nitrate, the sCOD utilized for denitrification would have been 70±90 mg/L in R1, 100±500 mg/L in R2, and 50±300 mg/L in R3. The percent sCOD utilized for denitrification during anaerobic influent distribution was 35% in R1, 75% in R2, and 37% in R3 of the total sCOD utilized during anaerobic influent distribution.
Figure 16 Nitrite/nitrate concentrations at the beginning and end of anaerobic influent distribution during steady-state for R1-R3. Error bars represent one standard deviation.

Characteristics and morphology of biomass with SRT

Steady-state MLVSS was 3.4±0.6 g/L in R1, 3.3±0.8 g/L in R2, and 1.8±0.2 g/L in R3. The difference in MLVSS between R1-R3 was determined to be significantly different (i.e., P <0.0001). Therefore, the OLR had a significant impact on steady-state MLVSS. The VSS/TSS of mixed liquor solids was determined to be 0.9±0.2 in R1, 0.9±0.3 in R2, and 0.9±0.2 in R3. Similarly, the VSS/TSS of effluent solids was determined to be 0.8±0.4, 0.8±0.5, and 0.9±0.5. The VSS/TSS of mixed liquor was not significantly different from that of the effluent solids. The actual SRT was estimated to be 7±2 d in R1, 7±3 d in R2, and 7±1 d in R3. The SRT$_{AGS}$ was estimated to be 12.5 d. Even at low SRT it was possible for AGS to remain stable and obtain >80% COD removal from the brewery wastewater.
Net observed yields were estimated to be 0.11±0.03 g-VSS/g-COD_{influent} for R1, 0.16±0.04 g-VSS/g-COD for R2, and 0.18±0.04 g-VSS/g-COD for R3 – Figure 17. The net observed yield increased as OLR decreased, suggesting that at high OLR (i.e., 4.5±0.6 kg-COD/(m³•d)) the conversion of substrate to new biomass was limited. The nitrogen content of MLVSS was determined to be 0.2±0.1 g-N/g-VSS in R1, 0.2±0.3 g-N/g-VSS in R2, and 0.2±0.2 g-N/g-VSS in R3. The nitrogen content of MLVSS was higher than what is typically observed for pure biomass, which can range from 0.07-0.12 g-N/g-VSS, values that are commonly used in conventional activated sludge models [21], [22]. The phosphorus content of MLVSS was determined to be 0.04±0.01 g-P/g-VSS in R1, 0.030±0.008 g-P/g-VSS in R2, and 0.023±0.005 g-P/g-VSS in R3. Typically, the phosphorus content of non-enhanced biological phosphorus removal biomass is 0.015 g-P/g-VSS [22].

The higher nitrogen content of AGS in this study may have been due to multiple factors, which cannot be concluded in this study. For instance, stable granular sludge has been observed to have higher protein to polysaccharide content [19], thereby potentially increasing the nitrogen content of solids. Furthermore, AGS has also been observed to act as an ion exchanger with ammonium and other ions in bulk solution [23]. In this study, due to the short SRT applied, it is possible that a higher quantity of nitrogen ions was being removed with the biomass during mixed liquor wasting. Similarly, the ion exchange mechanism would explain the higher phosphorus content of biomass, especially since enhanced biological phosphorus removal was not observed in this study. Like ammonium, phosphate ions have also been observed to accumulate in AGS [24].
Figure 17 Steady-state net observed yields shown with the nitrogen and phosphorus content of MLVSS. Error bars represent one standard deviation.

Based on the net observed yield, nitrogen content of MLVSS, and phosphorus content of MLVSS, it was possible to determine the minimum COD:TN:TP ratio for biomass growth. Previous studies have suggested a minimum ratio of 100 COD: 10 TN: 1 TP for brewery wastewater [25]. However, it was determined that a range of 100 COD: 2.4-2.8 TN: 0.4-0.5 TP would be enough for biomass growth in this study. Thus, nitrogen and phosphorus requirements based on the literature may have been overestimated by as much as 3.6-4.2 times and 2.0-2.5 times, respectively. This was a significant finding for high-strength wastewaters that have low nitrogen and phosphorus content, such as brewery wastewaters.
Impact of stopping SRT

After a month of steady-state operation, wastage of mixed liquor at an equivalent rate of 0.32 L/d was stopped. The AGS destabilized within several days, with what were originally large and uniform granules (i.e., from 0.5-5 mm) rapidly transitioning to particles less than 0.2 mm – Figure 18. Furthermore, the appearance of flocs and filamentous microorganisms was associated with the stoppage of SRT control. Without SRT control it was clear that granules in all three reactors, R1-R3, lost stability and rapidly disintegrated to smaller particles.

Destabilization of AGS, after mixed liquor wastage was stopped, was also associated with decreased performance in R1-R3 – Figure 19. Removal of COD and sCOD decreased from 81% to 51% and 78% to 62%, respectively, in R1. In R2 removal of COD decreased from 83% to 68% and sCOD removal decreased from 83% to 76%. COD and sCOD removal in R3 decreased from 86% to 76% and 95% to 92%, respectively. The associated RR for COD (i.e., \( R^2 = 0.9234 \)) and sCOD (i.e., \( R^2 = 0.9646 \)) correlated strongly with the OLR. The RRs for COD in R1-R3 were 0.63, 0.83, and 0.86, respectively. The RRs for sCOD in R1-R3 were 0.80, 0.91, and 0.97, respectively. Therefore, in terms of COD and sCOD removal, there was a higher impact on performance as OLR increased, suggesting that mixed liquor wasting was more important at higher OLR.
It was interesting to note that TN removal was improved five days after mixed liquor wasting was stopped. Total nitrogen removal in R1-R3 improved from 31%, 32%, and 33% to 36%, 45%, and 42%, respectively. This was likely due to the retention of more biomass, resulting in the accumulation of more nitrifying microorganisms with the flocculent and filamentous microorganisms that appeared. For instance, steady-state MLSS was 3.6 g/L in R1, 3.6 g/L in R2, and 1.9 g/L in R3. Five days after mixed liquor wasting was stopped, MLSS increased to 4.8 g/L in R1 (i.e., 130% increase), 6.6 g/L in R2 (i.e., 185% increase), and 6.9 g/L in R3 (i.e., 361% increase). As well, higher biomass concentrations may have also encouraged simultaneous nitrification-denitrification, although kinetic tests were not performed after mixed liquor wasting was stopped and therefore the onset of simultaneous nitrification-denitrification cannot be concluded.

It was also interesting to note that TP removal increased at the highest OLR (i.e., R1), while there was more TP in the effluent than in the influent at the lower OLRs (i.e., R2-R3). Specifically, TP removal during steady-state was 30% in R1, 35% in R2, and 34% in R3. Five days after mixed liquor wasting was stopped the TP removal in R1 was 43%, while it was -10% in R2 and -28% in R3. The reason for TP release in R2-R3 at the time of sampling may have been a result of phosphorus exchange between granules and solution, resulting in elevated phosphorus concentrations in bulk solution. Since the VER was higher in R1 (i.e., higher flow), the elevated phosphorus concentrations would have more quickly stabilized since the excess phosphorus would have been more quickly washed out of the system.
Figure 18 Stereoscopic images of biomass from R1-R3 during SRT control and seven days after SRT control was stopped.
Figure 19 Top – Difference in COD, sCOD, TN, and TP removals in R1-R3 with and without SRT control (i.e., mixed liquor wasting); Bottom – Removal ratios (RR) for COD, sCOD, TN, and TP as a function of OLR.
Summary

It was demonstrated the suitability of using SRT control to maintain AGS stability at high OLR and low hydrodynamic shear force (i.e., USAV = 0.51 cm/s). Granules were stable at OLR from 1.5-4.5 kg-COD/(m³•d) when SRT was controlled. It was determined that a range of 100 COD: 2.4-2.8 TN: 0.4-0.5 TP would be enough for biomass growth in this study. Granules quickly (i.e., within 5 d) lost their structural stability when mixed liquor wasting was stopped and SRT was not controlled. The RR for COD was 0.63 at 4.5 kg-COD/(m³•d), 0.83 at 3 kg-COD/(m³•d), and 0.86 at 1.5 kg-COD/(m³•d). The trend of RR with OLR suggested that mixed liquor wasting, and therefore SRT control, was more crucial at higher OLR. Thus, SRT control may be an adequate replacement for high hydrodynamic shear force to control AGS stability at high OLR.
Bibliography


[23] J. P. Bassin, M. Pronk, R. Kraan, R. Kleerebezem, and M. C. M. van Loosdrecht,


Chapter 5: Comparison of Hydrodynamic Shear Force and Granular Sludge Wasting as Methods to Stabilize AGS Biomass

Chapters 3 and 4 presented two methods for stabilizing AGS:

1. Hydrodynamic shear force; and
2. Controlled wasting of AGS.

Herein, the two methods for stabilizing AGS are described and discussed as they relate to reactor operation.

Hydrodynamic shear force

Chapter 3 examined the influence of increasing OLR on AGS-SBR operation under low hydrodynamic shear force. Specifically, USAV in the AGS-SBRs was 0.41 cm/s while OLR increased from 1.36 kg-COD/(m³•d) to 2.52 kg-COD/(m³•d) to 5.20 kg-COD/(m³•d). At 0.41 cm/s and 1.36 kg-COD/(m³•d), a mature AGS developed within 40 d. At OLR of 2.52 kg-COD/(m³•d) a mature AGS developed within a month, but filamentous microorganisms were detected on granule surfaces approximately 15 d after mature AGS development – Figure 20. Granules at OLR of 2.52 kg-COD/(m³•d) and USAV of 0.41 cm/s were fully coated by a filamentous surface layer within two months from inoculation. At OLR of 5.20 kg-COD/(m³•d) large granules developed rapidly, within two days, but then quickly destabilized. For the remaining four months of experimentation, biomass in the SBR with OLR of 5.20 kg-COD/(m³•d) and USAV of 0.41 cm/s appeared as filaments joined together by a white, viscous substance that was likely a mixture of EPS and some cells.
A review of the literature [1–5], combined with experimental results from Chapter 3, produced a consistent relationship between the OLR and USAV – Figure 21. From this relationship it was determined that approximately 0.2 cm/s of USAV was required per kg-COD/(m³•d) of OLR. At USAV higher than 0.2 cm/s per kg-COD/(m³•d), AGS was determined to be stable. However, any less than 0.2 cm/s per kg-COD/(m³•d) and AGS would risk the development of a filamentous surface layer or complete degranulation. This relationship provides a general estimate of aeration requirements to achieve stable granulation.
Figure 21 Cases of successful and unsuccessful granulation plotted with the OLR and USAV used in the respective study.

Based on 0.2 cm/s per kg-COD/(m$^3$•d), A$_{USAV}$ could be calculated (i.e., Appendix E).

Values for A$_{USAV}$ could then be compared to values for A$_{Nit}$ and A$_{Het}$. The ratio of A$_{USAV}$ to both A$_{Nit}$ and A$_{Het}$ (i.e., A/A) were then determined as a function of several operational and design parameters – Figure 22. When A/A was greater than 1, more air would be required to achieve the minimum USAV than the air supplied for wastewater treatment with or without nitrification. If A/A was less than 1, the airflow supplied for wastewater treatment with or without nitrification would provide USAV greater than the minimum required value. The results demonstrated that as SWL increased, values for A/A slightly increased as well. The results also demonstrated that as DO increased, values for A/A decreased significantly. It was also demonstrated that A/A was independent of OLR and reactor diameter. Thus, at lower SWL or higher DO, the air supplied for wastewater treatment with or without nitrification would approach or surpass the air required to achieve the minimum USAV. Dissolved oxygen in particular had the greatest impact on A/A.
It was also demonstrated that wastewater treatment with nitrification would approach an A/A of 1 sooner than wastewater treatment without nitrification. This was because nitrification requires additional oxygen to biologically oxidize ammonium to nitrate. Overall, modelled values for A/A demonstrated that AGS would be more suitable for treatment of wastewater that required nitrification. Therefore, AGS would be ideal for municipal wastewater treatment facilities where complete nutrient removal may be required. On the other hand, AGS may be less ideal for high-strength, organic wastewater with nitrogen deficiency (i.e., where nitrification would not be required). The reason being that additional aeration might be required to achieve the minimum USAV to guarantee AGS stability.
Controlled wasting of AGS

An alternative method for stabilizing AGS, compared to high hydrodynamic shear force, was examined in Chapter 4 while treating brewery wastewater. Specifically, the controlled wasting of AGS, in addition to the selective wasting of slowly-settling biomass, was examined at OLR from 1.5-4.5 kg-COD/(m$^3$•d) and low USAV of 0.51 cm/s. The measured whole sludge SRT was approximately 7 d under all OLR. Previously, in Chapter 3, a minimum USAV of 0.2 cm/s per kg-COD/(m$^3$•d) was recommended to maintain stable AGS. Thus, at 4.5 kg-COD/(m$^3$•d) the USAV should have been 0.9 cm/s to generate stable AGS. However, it was demonstrated that controlled wasting of AGS helped maintain stable AGS under lower USAV – Figure 23.

Therefore, controlled wasting of AGS could allow for lower USAV, thereby reducing aeration requirements. For municipal wastewater treatment, where nitrification was required, the USAV could be achieved by the airflow required for wastewater treatment. However, additional aeration would be required for purely heterotrophic processes, such as AGS for brewery wastewater treatment. Thus, controlled wasting of AGS would be an effective strategy for wastewaters with high organic content and low nitrogen content or for applications where nitrification was not required. Figure 24 presents a visual representation of granulation with increasing OLR with and without SRT control at low hydrodynamic shear force.
Figure 23 Successful granulation with SRT control shown with the OLR limit without SRT control and the potential OLR limit with SRT control. It should be noted that OLR above 4.5 kg-COD/(m³•d) were not examined, and therefore the OLR limit with SRT control may be higher than presented.

Figure 24 Visual representation of AGS stability with increasing OLR with and without SRT control at low hydrodynamic shear force.
Bibliography


Chapter 6: Description of the AGS-CFR

Introduction

AGS is an advanced biological process for energy efficient wastewater treatment in small footprints [1,2]. AGS offers several advantages over CAS including excellent settling, high biomass retention and therefore high volumetric loading, reduced oxygen demand, and lower sludge yields [3,4]. Furthermore, due to the dense and compact structure of granules, AGS offers higher resilience to drastic changes in operating conditions such as temperature, organic loads, and toxic compounds.

SBRs have been the technology of choice for most AGS studies, as well as for the only commercially available AGS process [5–7]. Sequencing batch reactors differ from most CAS processes, which use CFR, because SBRs combine all the treatment steps into a single basin. Instead of having multiple basins dedicated to different oxidation or reduction conditions, SBRs change operational parameters over time (e.g., anaerobic influent addition $\rightarrow$ aeration $\rightarrow$ settling $\rightarrow$ effluent discharge $\rightarrow$ repeat).

It was generally agreed upon that SBRs benefitted AGS and that granulation in CFRs would not be stable. The one distinct advantage SBRs have over CFRs is the ability to distribute influent through a settled bed of granules, thereby promoting anaerobic utilization of substrate. However, recent studies have demonstrated the potential for successful and stable granulation in CFRs [8]. These developments are important since most large WWTP practicing nutrient removal and requiring increased capacity are large CFRs, such as plug-flow reactors or completely-mixed reactors [9]. In these cases, existing infrastructure would not be suitable for retrofit to SBRs and it would be uneconomical to implement AGS unless a CFR was developed.
Accordingly, it was desirable to develop suitable process design and operational parameters for AGS-CFR, which could potentially enable high-levels of industrial and/or municipal wastewater treatment (i.e., carbon, nitrogen, phosphorus, and solids removal) with a smaller footprint and lower energy requirements than CAS processes.

Description of the reactor

The AGS-CFR comprises an anaerobic zone where influent is introduced to the AGS-CFR, an aerobic zone situated downstream of anaerobic zone, and an anoxic zone situated upstream of the anaerobic zone. A simplified schematic of the AGS-CFR is presented in Figure 25.

![Figure 25 Simplified schematic of the AGS-CFR in one of its embodiments.](Image)

Relative to the influent COD load, the total aerobic volume is preferably designed based on an OLR from 0.5-1.5 kg-COD/m³/d. The aerobic zone preferably includes at least one baffle or structure that provided adequate separation between the first 10-30% of the aerobic zone and the last 70-90%. The purpose of this separation would be to develop defined feast conditions (i.e., high substrate availability for growth) in the first, smaller compartment or subzone of the aerobic zone, while famine conditions (i.e., low substrate availability for growth) are developed in the second, larger compartment or subzone of the aerobic zone. The presence of defined feast and famine conditions, with approximately 20% or less, on average, of the aerobic volume
allocated for feast conditions, has been demonstrated to be beneficial for AGS stability as well as the selection of PAO over GAO [10,11]. Additional baffles or structures for separation could be applied in the aerobic zone to, likewise, create different compartments or subzones to facilitate control strategies for strict effluent nitrogen limits (i.e., TN <5 mg/L).

In the illustrated example, all wastewater influent is supplied through an inlet to the anaerobic zone, where readily available substrates are taken up by PAO and GAO, while more complex substrates would be broken down to less complex substrates [12]. The purpose of the anaerobic zone would be to encourage the development of PAO and GAO, reduce sludge yields by anaerobically degrading complex organic substrates, and direct carbon towards simultaneous nitrogen and phosphorus removing pathways. Wastewater with a readily bioavailable fraction of COD from 0.2-0.25 would be adequately treated with an anaerobic zone approximately 15-30% the volume of the aerobic zone. As the readily bioavailable fraction of COD increased, the volume of the anaerobic zone could be reduced. Likewise, as the readily bioavailable fraction of COD decreased the volume of the anaerobic zone may be increased to accommodate the degradation of a more complex wastewater substrate. Supplemental carbon may be required to treat wastewater with very low COD/TN.

A solids separation device, for example in the form of a secondary clarifier, resides downstream of the aerobic zone. A return line has an inlet connected to a sludge drain of the solids separation zone, and an outlet that opens into the anoxic zone at the upstream end of the reactor, whereby the return line recycles a stream of RAS from the solids separation zone back into the CFR. The anoxic zone would be designed based on endogenous respiration to completely remove all nitrites/nitrates in the sludge recycle. Operating the anoxic zone off endogenous material would act to extend famine conditions, with the intent of out-selecting
However, at lower sludge recycle rates, and thereby lower nitrate return, the anoxic zone may develop anaerobic conditions resulting in phosphorus release [13]. Therefore, the anoxic zone may be equipped with diffusers such that air may be supplied if anaerobic conditions are detected. This could be controlled by ORP.

As shown in Figure 25, WAS for selective wasting can be drawn from an elevated location inside the aerobic zone to a selective wasting device installed in operable relation thereto, while aerobic granular sludge can likewise be drawn from the aerobic zone from a lower elevation therein through a lower removal port communicating therewith. Alternatively, the selective wasting device and the removal port may be installed in cooperation with the return line to draw the waste activated sludge, both for selective wasting and wasting of aerobic granular sludge, from the return activated sludge RAS that is being recirculated through the return line. Similarly, other alternative locations for the selective wasting port include any elevation of the anoxic or anaerobic zones. A summary of design parameters for the AGS-CFR are presented in Table 8.

**Table 8 Summary of design parameters for the AGS-CFR.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Preferred range</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Configuration</td>
<td>anoxic/anaerobic/aerobic</td>
<td>SND in aerobic zone</td>
</tr>
<tr>
<td>Anoxic zone</td>
<td>single CMR/PFR or multiple CMR/PFR in series</td>
<td>Substrate gradients not as critical as aerobic zone</td>
</tr>
<tr>
<td>Anaerobic zone</td>
<td>single CMR/PFR or multiple CMR/PFR in series</td>
<td>Substrate gradients not as critical as aerobic zone</td>
</tr>
<tr>
<td>Aerobic zone</td>
<td>single PFR or multiple CMR/PFR in series</td>
<td>Cannot have a single CMR since feast and famine conditions will not develop</td>
</tr>
<tr>
<td>MLSS</td>
<td>4-12 g/L</td>
<td>Based on total reactor volume</td>
</tr>
<tr>
<td>OLR</td>
<td>0.5-1.5 kg-COD/(m³·d)</td>
<td>Relative to influent flow</td>
</tr>
<tr>
<td>Anoxic HRT</td>
<td>1-2.5 h</td>
<td>Relative to influent flow</td>
</tr>
<tr>
<td>Anaerobic HRT</td>
<td>1-3.5 h</td>
<td>Relative to influent flow</td>
</tr>
<tr>
<td>Aerobic HRT</td>
<td>4-14 h</td>
<td>Relative to influent flow</td>
</tr>
<tr>
<td>SRT&lt;sub&gt;AV&lt;/sub&gt;</td>
<td>0.5-6 d</td>
<td>SRT&lt;sub&gt;AV&lt;/sub&gt; may not be controlled during granulation (i.e., SRT&lt;sub&gt;AV&lt;/sub&gt; →∞)</td>
</tr>
<tr>
<td>SRT&lt;sub&gt;AGS&lt;/sub&gt;</td>
<td>30-50 d</td>
<td></td>
</tr>
<tr>
<td>R</td>
<td>0.2-1</td>
<td></td>
</tr>
</tbody>
</table>
Description of the process

Many individual control loops would contribute to the control system of a stable AGS-CFR. The first control system would monitor DO and other parameters in the aerobic zone. The simplest configuration, which would separate the aerobic zone into two discrete compartments or subzones (i.e., feast and famine), would control DO in the first aerobic compartment/subzone via a setpoint and DO in the second aerobic compartment/subzone via measured ammonium concentrations. A lower DO limit, such as 0.05-0.5 mg/L, would be applied in the second aerobic compartment/subzone to maintain an aerobic environment if the minimum ammonium concentration was achieved. In the case of more stringent effluent nitrogen limits (i.e., TN < 5 mg/L) and/or low carbon to nitrogen ratio in the influent, additional process controls could be added. Keeping with the simplest configuration (i.e., two aerobic compartments/subzones), online nitrite/nitrate monitoring could be implemented and linked to a carbon dosing system. If the total nitrite/nitrate concentration exceeded requirements, methanol or another carbon source providing low biomass yields could be dosed to the second aerobic compartment/subzone while the lower DO limit was operated (i.e., 0.05-0.5 mg/L).

In these cases, the volume of the second aerobic compartment/subzone would have to be extended to guarantee that ammonium would be within allowable concentrations while denitrification on supplemental carbon source was occurring. Otherwise, a more complex configuration with additional baffles or structures for flow separation within the second aerobic compartment/subzone (i.e., famine conditions), as well as additional probes for the separated compartments, could be employed with multiple carbon dosing locations. This would allow for optimal control of nitrification, denitrification, carbon dosing, and famine conditions, thereby
providing the most robust control system to meet low effluent nitrogen concentrations (i.e., TN of 3 mg/L).

The second control system, in its simplest form, would monitor ORP in the anoxic zone and turn aeration on if the ORP registered anaerobic conditions. The anoxic zone would serve two purposes, the first being a conventional purpose; preventing nitrite/nitrate carryover to the anaerobic zone. The second, novel purpose would be to serve as extended famine conditions for the AGS-CFR since the anoxic zone would be designed based on endogenous respiration. At low sludge recycle rates, especially while the AGS-CFR was under design capacity, the ORP may drop, resulting in anaerobic conditions. If this were to happen the anoxic zone would no longer serve as extended famine conditions (i.e., lack of electron acceptors) and secondary phosphorus release could occur [13]. Thus, it would be important to maintain ORP above anaerobic conditions, through aeration, to benefit system stability.

The second control system could be built upon through the addition of a nitrite/nitrate probe and carbon dosing system (i.e., methanol or equivalent). As the AGS CFR begins to reach capacity, or if influent carbon to nitrogen ratio was low, endogenous respiration may not be economical to completely reduce all nitrite/nitrate in the sludge recycle. Thus, a nitrite/nitrate probe could be used to monitor oxidized nitrogen species in the anoxic zone and turn on carbon dosing when nitrite/nitrate concentration exceeded desired values.
Description of potential modifications

In addition, or alternative to the above described features of the illustrated embodiment, the system and operational process can be modified in numerous ways. For example, an aeration grid can be added to all or part of the anoxic zone to make it an anoxic/aerobic swing zone capable of preventing deep anaerobic conditions from developing and thereby mitigating phosphate release that could be detrimental to biological phosphate removal [13,14]. Optionally operating the anoxic zone as a swing zone would also allow for extended starvation/famine conditions, which have been demonstrated to enhance granular sludge stability [10]. An aeration grid could also be added to all or part of the anaerobic zone to make it an anaerobic/aerobic swing zone capable. Operating all or part of the anaerobic zone as a swing zone would allow for the anaerobic HRT to be decreased, for example at lower substrate load and/or higher temperature, while still promoting the development of feast conditions.

A portion of the raw wastewater may be redirected from the main influent inlet through a diversionary intake branch, or a supply of carbon from an external carbon source can be introduced to the anoxic zone through an influent or carbon intake to accelerate denitrification rates [15,16] when the nitrite/nitrate load in the return sludge stream is higher than the design load. Likewise, a portion of the raw wastewater may be redirected from the main influent through a divisionary intake branch, or a supply of carbon from an external carbon source can be introduced to the anaerobic zone in multiple anaerobic compartments, whether multiple CMRs in series or different locations along a PFR, to distribute the carbon source, internal or external, throughout the anaerobic zone.
A portion of the raw wastewater may also be redirected from the main influent through a
divisionary intake branch, or a supply of carbon from an external carbon source can be
introduced to the portion of the aerobic zone, whether multiple CMRs in series or different
locations along a PFR, that is allocated for feast conditions. The portion of carbon, internal or
external, diverted to the aerobic zone allocated for feast conditions would maintain feast
conditions during periods of concern, such as low OLR. With regards to solids inventory, a
portion of the return activated sludge may be redirected from the main return activated sludge
inlet to the anoxic zone through a diversionary branch to an inlet to the aerobic zone. The
diversion of a portion of the return activated sludge to the aerobic zone could be utilized to
extend famine conditions and minimize the nitrite/nitrate load to the anoxic zone at higher return
activated sludge flows.
Bibliography


[14] P. Jabari, Q. Yuan, J.A. Oleszkiewicz, Overall effect of carbon production and nutrient release in sludge holding tank on mainstream biological nutrient removal efficiency,

Chapter 7: Influence of selective pressure in an AGS-CFR

Introduction

One of the key factors for successful and stable granulation, in SBR or CFR, is the cultivation of beneficial microorganisms such as GAO and PAO [1], [2]. Accumulating microorganisms allow for uniform utilization of organic substrate throughout granule depth, thereby redirecting substrate from fast-growing aerobic microorganisms [3], [4]. Furthermore, dense and rapidly settling granules have been observed to contain a higher proportion of PAO than GAO [5], [6]. This is likely due to high concentrations of polyphosphates and other inorganics in PAO dominated granules [7]. For these reasons, AGS-CFRs must consider microbial selection pressures that favour the growth of PAO to achieve stable and rapidly settling granules. Such considerations include dedicated anaerobic zones where readily available substrate is utilized by accumulating microorganisms as well as selective wasting that separates the light fractions of biomass from rapidly settling PAO aggregates.

It has also been demonstrated that extended famine conditions promote AGS stability [8]. Phosphorus accumulating organisms out-compete GAO at low organic loadings rates, or extended periods of aerobic decay, known as famine conditions [9]. This was because PAO tend to maintain a higher reserve of polyhydroxyalkanoate than GAO, thereby providing PAO with an energy source for aerobic maintenance processes. Therefore, AGS-CFRs would benefit from extended aerobic or anoxic conditions with regards to selection of PAO over GAO.

Hydrodynamic shear force was also suggested to influence granule stability, although that has been refuted under low organic loading [3], [10]. Existing full-scale AGS-SBRs are operated at OLR near 1 kg-COD/(m³•d) [11], although stable AGS under high hydrodynamic shear force have been reported at organic loading rates up to 15 kg-COD/(m³•d) [12]. For municipal
wastewaters, which are typically low-strength, AGS-CFRs would not require high hydrodynamic shear force. However, industrial wastewaters with high bCOD loads (e.g., brewery wastewater [13]) may benefit from higher hydrodynamic shear force.

One of the first studies on an AGS-CFR demonstrated that the granular inoculum rapidly lost its structural integrity, resulting in loose and fluffy aggregates, after being transferred to the CFR [14]. A more recent study examined an AGS-CFR with two settling tanks in series, although micropowder addition was required to enhance granulation [15]. Otherwise, there is a distinct lack of knowledge in the literature about AGS-CFRs. The objective of this study was to determine the impact of selective pressure, via selective wasting, on granulation of flocculent biomass in a CFR designed for PAO cultivation. The goal was to demonstrate that aerobic granulation was possible in a CFR, and that a biomass with high treatment capacity (i.e., HRT of 6 h) could be developed.

Methods

*Reactor configuration*

A modular CFR was constructed from HDPE containers of various working volumes (i.e., 5-40 L) in series and operated at 20±1 °C. Each HDPE container represented a single CMR and was mixed with 5” (i.e., 12.7 cm) right-handed propellers rotating clockwise. Rotational speed of the propellers was adjusted to ensure adequate mixing in each CMR (i.e., 15 rpm for 5 L; 120 rpm for 40 L). A series of CMRs was configured using through-wall fittings centered 1” (i.e., 2.54 cm) above the bottom of the container. Containers were connected with food-grade plastic tubing with an inside diameter of 3/8” (i.e., 0.9525 cm).
Wastewater was supplied with peristaltic pumps at a total flow of 240 L/d and the HRT was 6 h. The flow was distributed equally between the first three 5 L reactors (i.e., 80 L/d to each reactor). The first 5 L reactor, which also received the sludge recycle (i.e., constant at 168 L/d which was 70% total influent flow of 240 L/d) acted as the pre-anoxic zone while the following two 5 L reactors operated as anaerobic zones. This “step-feed” approach was intended to maximize the anaerobic HRT while mitigating issues associated with extended anaerobic conditions in the presence of low substrate concentrations. Dissolved oxygen was monitored with optical probes in both the 5 L and 40 L aerobic zones. A programmable logic controller received input from the optical probes and relayed information to outlets controlling air pumps on/off. Dissolved oxygen was maintained between 1.5-2.5 mg/L in both the 5 L and 40 L aerobic tanks. The aerobic zone was split into a small (i.e., 5 L) and large (i.e., 40 L) compartment in order to balance a short feast period in the first, small compartment followed by a long famine period in the second, large compartment [8].

To simplify analysis and operation of the lab-scale AGS-CFR, selective wasting was coupled with solid-liquid separation in the 5 L clarifier which had a surface area of ~573 cm². Scale-up of the process would require a discrete selective wasting device that diverted selectively wasted solids away from the final effluent. Such a device could draw biomass for selective wasting from any point in the reactor configuration. In the lab-scale AGS-CFR, selective wasting was achieved with a 5” (i.e., 12.7 cm) right-handed propeller rotating clockwise in the clarifier. Higher rotational speeds, with a maximum of 25 rpm, resulted in more turbulence during settling, and therefore higher selective pressures for rapidly settling particles.

The shear force applied during selective wasting, or mixing in general, was not expected to influence granulation since previous studies has demonstrated that high shear forces are not a
requirement for granulation [3]. Furthermore, the confounding influence of shear force applied
during selective wasting should have been minimized due to the small range of rotational speeds
examined (i.e., 7-25 rpm) for selective wasting, as well as due to much higher rotational speeds
(i.e., up to 120 rpm) being applied in the reactor compartments for mixing. Four distinct values
of selective wasting pressure were applied (i.e., 25, 20, 15, and 7 rpm). The first selective
wasting pressure examined was the maximum of 25 rpm since higher degrees of wasting would
have resulted in lower whole sludge SRT and therefore steady-state would have been attained
more rapidly. The CFR was disinfected and inoculated with flocculent biomass before a different
selective pressure was applied.

Source of inoculum

The AGS-CFR was inoculated with conventional activated sludge from a full-scale
biological nutrient removal municipal wastewater treatment plant located in Winnipeg, Canada.
The full-scale process operates in a Westbank configuration with organic matter oxidation,
nitrification of ammonium, denitrification for TN removal, and enhanced biological phosphorus
removal achieving effluent quality of <15 mg-TN/L and <1 mg-TP/L. Sludge volume index
typically varies from 120-160 mL/g. The full-scale process also utilizes primary clarifiers to
remove particulate organic matter prior to biological treatment. The primary solids from
clarification are fermented to yield volatile fatty acids, which are recycled to the anaerobic zone
of the Westbank process to assist the enhanced biological phosphorus removal. The inoculum,
which was RAS from the full-scale Westbank reactor, was introduced such that the original TSS
concentrations in the AGS-CFR was approximately 2 g/L. Whenever a new selective pressure in
the AGS-CFR was examined (i.e., rotational speeds of 7-25 rpm), the previous biomass was
removed, and the process was re-seeded with fresh biomass from the full-scale Westbank reactor.

**Wastewater characteristics**

Wastewater was composed of beef and yeast extracts (Sigma Aldrich), ammonium chloride, and potassium phosphate dibasic. Wastewater characteristics were determined on samples from the inlet of the AGS-CFR. On average COD was 370±30 mg/L, TN was 43±7 mg/L, and TP was 10±2 mg/L. The fraction of COD that was soluble was 0.89, while the fraction of TN that was TAN was 0.44 and the fraction of TP that was OP was 0.70. Based on steady-state data from operation with the least amount of selective pressure applied, and therefore the highest concentration of biomass, the fraction of nbsCOD in the wastewater was estimated to be 0.07±0.02. The ratio of COD to TN and COD to TP in the wastewater were 8.7±0.2 and 35.8±0.2, respectively. Approximately 220±50 mg/L of COD would be required to achieve effluent concentrations of 15 mg-TIN/L and 1 mg-OP/L (i.e., assuming NOx-N = 0.8 TN, 10 g-bCOD/g-P, and 6.6 g-bCOD/g-N).

**Analytical methods**

Influent and effluent samples were analyzed for COD, sCOD, TN, and TP spectrophotometrically (TNTplus, Hach, CA). TAN, nitrite nitrogen, nitrate nitrogen, and OP were also quantified spectrophotometrically (Lachat QuikChem 8500, HACH, CA). Samples for MLSS, MLVSS, and effluent TSS and VSS were analyzed according to Standard Methods [16]. A SteREO Discovery (Zeiss, DE) was used for microscopic analysis. All samples that required filtration were run through medium porosity Q5 filter paper (Fisher Scientific, CA). ImageJ was utilized to assess DF and DS. A minimum cutoff of 0.2 mm on DF was used to assess average diameter of granules [4]. The ratio of DF to DS therefore represented uniformity of granular
sludge particles, with higher ratios representing more irregular granular sludge particles. One-way analysis of variance (VassarStats, USA) was used to determine if steady-state data at different selective wasting pressures was significantly different (i.e., α = 0.05). \( Y_{\text{net}} \) was estimated according to Equation 5 and anaerobic sCOD utilization was estimated according to Equation 14.

\[
\text{Anaerobic sCOD utilization} = \left( 1 - \frac{sCOD_{\text{last anaerobic zone}} \cdot Q_{\text{influent}} \cdot (1 + R)}{sCOD_{\text{effluent}} \cdot Q_{\text{influent}} \cdot R + sCOD_{\text{influent}} \cdot Q_{\text{influent}}} \right) \cdot 100
\]

Results

**Biomass quantities and characteristics**

The quantities and characteristics of mixed liquor in the AGS-CFR were determined during steady-state operation at different selective wasting pressures – Figure 26. The degree of selective wasting applied, represented by rotational speeds of the impeller from 7-25 rpm in the clarifier, had a significant impact on MLSS (i.e., \( P < \alpha = 0.05 \)). At higher rotational speeds the steady-state MLSS was lower, approaching 150 mg/L. This could be explained by higher degrees of biomass washout as rotational speed increased, since the critical settling velocity in the clarifier would have been higher. At the lowest rotational speed examined, which was 7 rpm, the steady-state MLSS was observed to be >3000 mg/L. Furthermore, an impact on the VSS/TSS ratio of the mixed liquor was also observed with selective wasting. In general, the VSS/TSS ratio of the mixed liquor increased from 0.71 at a rotational speed of 15 rpm to 0.93 at a rotational speed of 25 rpm.
Figure 26 Steady-state MLSS and the ratio of MLVSS/MLSS with increasing selective pressure (i.e., rotational speed of the impeller in the clarifier). Error bars represent a single standard deviation.

Since the higher rotational speeds would have resulted in higher degrees of biomass washout, the accumulation of inorganic material in the mixed liquor would not have been as significant when compared to lower degrees of biomass washout. For this reason, lower rotational speeds would have resulted in the retention of more biomass and therefore allowed for the accumulation of more inorganic material. In general, it has been observed that stable AGS systems accumulated inorganic phosphate precipitates, such as calcium phosphate, in the granule core [7]. Thus, in addition to having excellent effluent characteristics under the steady-state sampling periods, rotational speeds ≤15 rpm also resulted in the accumulation of more inorganic matter (i.e., lower VSS/TSS in the mixed liquor), suggesting greater potential for stable granulation.
Based on the clarifier dimensions and impeller properties, the applied rotational speed was converted to a Re and velocity gradient. At the lowest rotational speed applied of 7 rpm the Re was 1434 and the estimated velocity gradient was 1 1/s. At the highest rotational speed applied of 25 rpm the Re was 5120 and the estimated velocity gradient was 9 1/s. At Re from 1434-5120, the flow in the clarifier was neither laminar nor turbulent and existed in-between those ranges. The Re and/or velocity gradient imparted by the impeller on the fluid in the clarifier could be used as a translatable measure for upscaling the selective wasting process. Based on observed steady-state MLSS, Re between 3000-4000 (i.e., 15-20 rpm), or velocity gradients between 4-7 1/s, had the greatest influence on biomass quantities.

Morphology of the biomass was also significantly impacted by the applied selective pressure. At all applied selective pressures there was a persistent flocculent fraction, which was visibly the highest at 7 rpm. The largest granules were observed at >20 rpm, although a significant fraction of filamentous microorganisms was visible at 25 rpm. Filaments were also present at 20 rpm where they were mostly associated with the surface of granular sludge particles. This was different than at 25 rpm where the filaments were also free-living and separated from granular sludge particles. Granular sludge was observed at all rotational speeds, although granules developed at 7 rpm did not meet the 0.2 mm minimum Df cutoff for measurement – Figure 27.
At a rotational speed of 15 rpm granules had an average $D_F$ of 0.22±0.01 mm and average $D_S$ of 0.11±0.02 mm. There was a visual reduction in the flocculent biomass fraction at 15 rpm compared to 7 rpm. At 20 rpm the average $D_F$ and $D_S$ were 0.4±0.2 mm and 0.3±0.1 mm, respectively. Similarly, at 25 rpm the average $D_F$ and $D_S$ were 0.31±0.09 mm and 0.18±0.08 mm, respectively. The ratio of $D_F$ to $D_S$ increased as rotational speed of selective wasting increased, suggesting that granules were more irregularly shaped at higher selective wasting pressures. The ratio of $D_F$ to $D_S$ was 1.3, 2.4, and 3.3 at rotational speeds of 15 rpm, 20 rpm, and 25 rpm, respectively. Overall, these results indicated that granulation under continuous flow is possible, and that significant performance can be achieved with the appropriate applied selective pressure.

Figure 27 Average $D_F$ and $D_S$ of granules in the AGS-CFR with increasing selective pressure. Error bars represent a single standard deviation.
Effluent characteristics

Steady-state effluent characteristics were quantified for the AGS-CFR at each of the selective wasting pressures applied (i.e., rpm of 7, 15, 20, and 25 in the clarifier) – Figure 28. Effluent TSS, VSS, COD, TN, and TP were not considered as performance measures for effluent characteristics since selective wasting was performed with solids liquid separation, which simplified lab-scale operation. Selectively wasted solids would have been present in the effluent and therefore confounded performance measures for effluent characteristics. For this reason, performance measures for effluent characteristics were sCOD, TIN, TAN, and OP since they are not influenced by effluent TSS and VSS concentrations.

Figure 28 Steady-state effluent characteristics for the AGS CFR under different selective wasting pressures. Higher selective wasting pressures were applied at higher rotational speeds of the impeller in the clarifier. Error bars represent one standard deviation.
In general, the performance of the AGS-CFR improved as the rotational speed decreased from 25 rpm to 7 rpm. The difference in sCOD, TIN, TAN, and OP between rotational speeds of 7-15 rpm were not significant (i.e., \( P > \alpha = 0.05 \)). However, a significant difference in effluent characteristics was observed for rotational speeds \( \geq 20 \) rpm and those \( \leq 15 \) rpm. Specifically, at rotational speeds \( \geq 20 \) rpm the steady-state sCOD was \( \geq 90 \pm 40 \text{ mg/L} \), TIN was \( \geq 18 \pm 2 \text{ mg/L} \) and OP was \( \geq 4 \pm 3 \text{ mg/L} \). On the other hand, below 15 rpm the steady-state sCOD was \( \leq 50 \pm 30 \text{ mg/L} \), TIN was \( \leq 17 \pm 8 \text{ mg/L} \) and OP was \( \leq 0.3 \pm 0.2 \text{ mg/L} \). Based on the observed effluent characteristics for rotational speeds \( \leq 15 \) rpm, which were determined from a month of steady-state operating data, the AGS-CFR would be capable of meeting stringent effluent characteristics. In fact, at 7 rpm effluent sCOD was \( 25 \pm 7 \text{ mg/L} \), TIN was \( 11 \pm 1 \text{ mg/L} \), and OP was \( 0.1 \pm 0.1 \text{ mg/L} \).

These results demonstrated that AGS can be successfully cultivated in continuous flow from a flocculent inoculum and achieve simultaneous COD, TN, and TP removal at a low hydraulic retention time of 6 h. These results also suggested that conversion of full-scale continuous flow facilities could be achieved by gradually increasing the selective pressure, allowing for continuous performance while transitioning from a flocculent to granular sludge process.

**Metabolic characteristics and biomass yields**

The first aspect of metabolism considered for assessing stable granulation was the development of distinct feast and famine phases. Previous research has demonstrated that a famine period of at least 80% the aerobic HRT cultivates a more stable aerobic granular sludge (Corsino et al., 2016). The AGS-CFR examined in this study contained two aerobic compartments with a total volume of 50 L. The first aerobic compartment, where feast conditions would have developed, was sized at 5 L. Thus, the second aerobic compartment, which was 40
L, provided 90% of the total aerobic volume for famine conditions to develop. However, it was observed that the actual development of famine conditions depended on the applied selective wasting pressure – Figure 29.

![Figure 29 Steady-state concentrations of sCOD in the first and second aerobic compartment of the AGS-CFR under different rotational speeds for selective wasting. Error bars represent one standard deviation.](image)

Steady-state sCOD concentrations in the first and second aerobic compartment provided quantifiable measures of feast and famine periods in the AGS-CFR. Thus, when sCOD in the first aerobic compartment was similar to sCOD in the second aerobic compartment (i.e., ratio of sCOD in first aerobic compartment to second aerobic compartment approaching 1) famine conditions would have developed. This was the case for rotational speeds of 7-15 rpm, with ratios of sCOD in the first to second aerobic compartments from 0.7±0.4-1.1±0.2. However, when the rotational speed was ≥20 rpm, the ratio of sCOD in the first to second aerobic compartment was greater than 1.6±1.
Based on measured sCOD concentrations, the second aerobic compartment was oxidizing 63-70 mg/L of sCOD on average, or 15.1-16.8 g/d of sCOD. Normalized to the 40 L volume of the second aerobic compartment, volumetric sCOD utilization rates were anywhere from 0.34-0.37 g/(L·d) for rotational speeds greater than 20 rpm. On the other hand, there was insignificant oxidation of sCOD for rotational speeds of 15 rpm and below, demonstrating the development of famine conditions in the second aerobic compartment. Thus, selective wasting pressures imparted by rotational speeds of 20 rpm and above resulted in steady-state MLSS concentrations that were not able to fully utilize the sCOD load in the first aerobic compartment. The lack of sufficient biomass subsequently resulted in a breakthrough of sCOD to the second aerobic compartment and a persistent feast phase that is associated with unstable granulation.

The second metabolic aspect considered to assess the stability of granular sludge was the degree of anaerobic substrate utilization – Figure 30. It is generally agreed that higher degrees of anaerobic substrate utilization result in more stable granulation due to reduced biomass yields and a uniform growth of microorganisms across granule depth [4]. At rotational speeds of 20 rpm and above the amount of anaerobic sCOD utilized was insignificant. Specifically, only 9±4% of sCOD was anaerobically utilized at 20 rpm while 10±10% was anaerobically utilized at 25 rpm. Steady-state values for anaerobic utilization of sCOD were much higher at rotational speeds of 15 rpm and lower. For instance, the AGS-CFR was able to utilize 45±3% of sCOD anaerobically at a rpm of 15. At a rpm of 7 the AGS-CFR anaerobically utilized 59±2% of the sCOD. Thus, in addition to disrupting adequate feast and famine regimes, rotational speeds of 20 rpm and above resulted in poor anaerobic utilization of sCOD, which could result in poor granular sludge stability.
The reason that overall anaerobic sCOD utilization was lower at higher rpm is because of the lower MLSS observed with increasing rotational speed of selective wasting. Higher rotational speeds of selective wasting resulted in more aggressive biomass washout and lower steady-state concentrations of MLSS. Thus, there was inadequate biomass quantity available to contact and uptake the entire sCOD load during anaerobic conditions. However, it is interesting to note that as rotational speed of selective wasting increased the specific anaerobic sCOD utilization rate also increased. For instance, at 7 rpm the specific anaerobic sCOD utilization rate was 16.7±0.5 mg-sCOD/(g-VSS·min), while at 15 rpm, 20 rpm, and 25 rpm the specific anaerobic sCOD utilization rates were 16.9±0.8 mg-sCOD/(g-VSS·min), 60±20 mg-sCOD/(g-VSS·min), and 70±70 mg-sCOD/(g-VSS·min), respectively.
Based on these observations it can be concluded that higher selective wasting pressures resulted in a biomass with higher anaerobic sCOD utilization rates. Therefore, if the steady-state MLSS concentrations had been higher at >20 rpm, the overall anaerobic sCOD utilization would have been improved. However, excessive washout of biomass at >20 rpm alluded that optimal granulation would have occurred somewhere between 15 rpm (i.e., high overall anaerobic sCOD utilization) and 20 rpm (i.e., high specific anaerobic sCOD utilization) in the AGS-CFR examined.

Observed yields for the AGS-CFR were determined based on the total quantity of wasted biomass and amount of COD removed – Figure 31. The simplified nature of the AGS-CFR meant that solids in the effluent were equal to the total quantity of biomass wasted (i.e., selective wasting was performed in the clarifier). Effluent VSS was lowest, at 23 mg/L, when rotational speed was 15 rpm. Highest effluent VSS of 130 mg/L was generated by the highest rotational speed of 25 rpm. This can be explained by higher critical settling velocities in the clarifier with higher rotational velocities, thereby resulting in more biomass washout and the development of a biomass with high-rate proprieties such as higher observed yields. However, at a rotational speed of 7 rpm, effluent VSS of 70 mg/L was higher than effluent VSS at 15 rpm. Likewise, observed yield increased from 0.08-0.44 g-VSS/g-COD as rotational speed increased from 15-25 rpm. Interestingly, at a rotational speed of 7 rpm the observed yield of 0.19 g-VSS/g-COD was higher than at 15 rpm.
Whole sludge SRT (i.e., considering total biomass and not the fraction of granules or flocs) was also calculated from steady-state parameters – Figure 32. At rotational speeds of 20 rpm and above, the SRT was 6.5-10 h, which would be an extremely aggressive SRT representative of a high-rate aerobic metabolism. At a rotational speed of 7 rpm the whole sludge SRT was 8.6 d, while the highest whole sludge SRT of 18 d was observed at a rotational speed of 15 rpm. Under the conditions examined a rotational speed of 15 rpm provided optimal sludge wasting and selected dense aggregates with good metabolic capabilities (i.e., not high-rate aerobic metabolism with high observed yields) while preventing accumulation of the flocculent, light fraction of biomass. At a rotational speed of 7 rpm the selective pressure was not significant enough to remove the flocculent, light fraction of biomass. This allowed for the accumulation of additional flocculent solids in the mixed liquor.

Figure 31 $Y_{net}$ under different rotational speeds for selective wasting.
Since there was no other method of biomass wasting, other than with the effluent, the flocculent, light fraction of biomass grew to the conditions provided at 7 rpm (i.e., biomass grew until clarification capacity was not sufficient to handle the solids loading rate). This explained why the effluent VSS and observed yield was higher at a rotational speed of 7 rpm than at a rotational speed of 15 rpm. Out of the four rotational speeds examined in this study, 15 rpm provided the optimal degree of selective wasting by selecting for granular sludge, minimizing the flocculent, light fraction of biomass, and preventing the development of high-rate conditions.
Summary

Formation of aerobic granular sludge was examined in a novel continuous flow configuration at 20±1 °C. Synthetic proteinaceous wastewater with municipal primary effluent characteristics was used (i.e., COD = 370±30 mg/L; TN = 43±7 mg/L; and TP = 10±2 mg/L). Various levels of selective pressure were applied after inoculation with flocculent sludge (i.e., estimated velocity gradients during settling between 1-9 1/s). Impeller rpm of 15 and below generated flocculent-granular biomass, while 20 rpm and above generated large granules with a filamentous population. Effluent soluble COD, TIN, and OP of 25±7 mg/L, 11±1 mg/L, and 0.1±0.1 mg/L, respectively, were obtained. Observed yields were as low as 0.08-0.19 g-VSS/g-COD and whole sludge SRT was 18±1 d. Famine conditions developed for 90% of the total aerobic volume and >45±3% anaerobic substrate utilization was recorded. Aerobic granulation was demonstrated feasible under continuous flow providing adequate treatment with low biomass yields.
Bibliography


Chapter 8: Impact of feast to famine ratios in an AGS-CFR

Introduction

The benefits of AGS cannot be realized by all treatment facilities since SBRs are not suitable for retrofitting continuous flow reactors that employ long, plug-flow concrete basins for biological nutrient removal. Thus, there is a demonstrated need to assess AGS-SBR design parameters and translate them to CFRs. Continuous flow reactors for AGS have seen moderate success in lab-scale configurations. Aerobic granular sludge was successfully cultivated in a CFR with two settling tanks in series, although micropowder addition was required to enhance granulation [1]. Another study demonstrated that granular sludge inoculum rapidly lost its structural integrity, resulting in loose and fluffy aggregates, when transferred to a membrane bioreactor [2].

Granulation was successful in approximated plug-flow with secondary clarification on synthetic low-strength wastewater [3]. In this case, granules were stable when famine conditions developed over 90% of the total aerobic volume, although granules became large and fluffy when feast conditions developed over 100% of the total aerobic volume. “Feast” conditions refer to the time during aeration when substrate, typically organic based, in bulk solution is high while “famine” conditions refer to the time during aeration when substrate in bulk solution is low [4]. Multiple AGS-SBR studies have also observed unstable granules when feast conditions are not properly balanced by adequate famine conditions, thereby suggesting that properly balanced feast and famine conditions are therefore important for both AGS-SBR and AGS-CFR applications [5], [6].
Specifically, extended famine conditions (i.e., a lower ratio of feast to famine duration) seemed to be important for the selection of PAO. Enrichment of PAO is important because dense granules have been observed to have a higher proportion of *Candidatus Accumulibacter phosphatis*; a commonly detected PAO [7]. Higher proportions of PAO have also been linked to higher inorganic content of granules, which leads to increased settling velocity [8], [9]. During famine conditions, PAO have been observed to maintain a higher reserve of PHA than GAO, and thereby PAO out-competed GAO during extended famine conditions [10]. Furthermore, manual removal of bulk substrate from solution before aeration (i.e., reduction of feast conditions) resulted in increased PHA storage capacity of biomass; from 48% up to 70% of total biomass weight [11]. Increased PHA content would be related to a higher quantity of PAO and/or GAO, therefore adequate balancing of feast and famine conditions can lead to the enrichment of PAO and better performing AGS.

The objectives of this study were to demonstrate the potential for granulation in a pilot-scale CFR, without selective wasting, while treating low-strength, real domestic wastewater. The impact of reactor of configuration, specifically the volume ratios of the first to second aerobic zone, on AGS formation and stability was examined. A larger ratio of the first to second aerobic zone represented longer feast conditions, while a smaller ratio represented longer famine conditions. Selective wasting was then incorporated to the optimal reactor configuration to examine the development of mature AGS under optimal conditions.
Methods

Reactor configuration and operation

A modular CFR was constructed out of multiple completely mixed reactors in series. Thus, the CFR configuration could be changed by adding or removing individual completely mixed reactors. The modular CFR was constructed from plastic tanks of various working volumes (i.e., 30-300 L) in series. The tanks were connected via ¾” (i.e., ~1.91 cm) inside diameter plastic tubing and flow through the CFR was driven by gravity. Wastewater from an equalization tank was pumped into the first anaerobic zone. RAS was also pumped to the first anaerobic zone from the pre-anoxic zone. The zones were aligned such that mixed liquor flowed from the first anaerobic tank and through the second anaerobic tank, first aerobic tank, second aerobic tank, and clarifier. Clarified effluent was discharged via overflow from the clarifier weirs and RAS was conveyed by gravity to the pre-anoxic zone.

In configuration 1 the total volume was 740 L with a total anoxic volume of 100 L (i.e., 13.5% total volume), a total anaerobic volume of 200 L (i.e., 27% total volume), and a total aerobic volume of 440 L (i.e., 59.5% total volume). In configuration 2 the total volume was 650 L with a total anoxic volume of 120 L (i.e., 18.5% total volume), a total anaerobic volume of 200 L (i.e., 31% total volume), and a total aerobic volume of 330 L (i.e., 50.5% total volume). The target HRT was 12 h for each configuration. Thus, the target anaerobic and aerobic HRTs in configuration 1 were 3.2 h and 7 h, respectively. In configuration 2 the target anaerobic and aerobic HRTs were 3.7 h and 6 h, respectively. The clarifier had a fixed volume of 100 L and HRT of 1.6 h in configuration 1, and HRT of 1.8 h in configuration 2.
A 120 L sidestream selective wasting device was added to configuration 2, which was determined to be the best performing configuration. The sidestream selective wasting device drew mixed liquor from the 300 L aerobic tank and subjected biomass to a settling regime before discharging the light fraction of biomass. The settled biomass in the sidestream selective wasting device was resuspended and returned to the 300 L aerobic tank. The sidestream selective wasting device operated on a 4 h cycle with 208-212 min aeration, 8-12 min settling, 10 min discharge of selective waste, and 10 min idling. During aeration biomass was continuously pumped from the 300 L aerobic tank of configuration 2 to the sidestream selective wasting device. An overflow from the sidestream selective wasting device, located at a height of 50 cm, allowed for mixed liquor to be returned to the 300 L aerobic zone. The pump ran at 1.5 L/min, thereby allowing for 3 volume exchanges of the sidestream selective wasting device before each settling period. The discharge port for selective waste was located at a height of 25 cm, thus the biomass had to travel 25 cm (i.e., 50 cm – 25 cm) within the allocated settling time to be retained in the reactor. At settling times of 12, 10, and 8 min the respective critical settling velocities were approximately 2.1, 2.5, and 3.1 cm/min.

Each reactor tank was completely mixed with 5” (i.e., 12.7 cm) right-handed propellers rotated clockwise by mechanical mixers. Aeration grids in the first and second aerobic zone were constructed from ½” (i.e., ~0.197 cm) copper tubing with holes spaced every 2” (i.e., ~0.787 cm). Air was continuously supplied to the first aerobic zone by a blower and DO in the first aerobic zone ranged from 2-4 mg/L. Air was intermittently supplied to the second aerobic zone by a blower turned on at a lower DO limit of 0.5 mg/L and an upper DO limit of 2 mg/L (i.e., IQ SensorNet 2020, Avensys Solutions Inc., CA). The clarifier had a scum scraper rotating at 3 rpm to prevent accumulation of float solids. In both configurations, mixed liquor was wasted by
pumping from the second aerobic zone for SRT control. The target SRT, by wasting, in both cases was 40 d. The actual SRT was lower than the target SRT in both cases because of effluent solids (i.e., an additional point of wasting). The recycle ratio, R, of RAS was targeted to be 1 (i.e., 100% of influent flow).

*Source of inoculum*

The AGS-CFR was inoculated with conventional activated sludge from a full-scale biological nutrient removal domestic wastewater treatment plant located in Winnipeg, Canada. The full-scale process operates a Westbank-type configuration with organic matter oxidation, nitrification of ammonium, denitrification for TN removal, and enhanced biological phosphorus removal achieving effluent quality of <15 mg-TN/L and <1 mg-TP/L. The process utilizes primary clarifiers for the capture of particulate organic matter prior to biological treatment. Primary solids are fermented at the full-scale plant to yield volatile fatty acids, which are conveyed to the anaerobic zone of the Westbank-type process to assist enhanced biological phosphorus removal. It should be noted that the pilot AGS-CFR operated on primary effluent without the addition of volatile fatty acids from primary solids fermentation. The inoculum was introduced such that the original TSS concentrations in the AGS-CFR was at least 2 g/L, which was deemed an adequate concentration for lab-scale AGS-CFR studies [3].
During maintenance periods in the full-scale plant (e.g., replacement of fine bubble air diffusers) 35 mg/L of ferric chloride was added to the raw wastewater after bar screening for chemically enhanced primary treatment. It should be noted that ferric chloride was added during start-up of the AGS-CFR (i.e., days 40-100). However, the AGS-CFR was re-seeded with inoculum before transition to configuration 1 and again re-seeded before transition to configuration 2. An acclimation phase of 30-40 d after inoculation was allowed before steady state analysis was considered.

Wastewater characteristics

Effluent from the full-scale plant’s primary clarifiers was supplied to the AGS-CFR. The primary effluent was continuously gravity-fed to a 600 L equalization tank with hydraulic circulation/mixing via a submersible pump. To mitigate fermentation the equalization tank was emptied and cleaned 2-3 d per week. The equalization time provided was 2.5-6.5 h depending on how much flow was being drawn for the AGS-CFR and other pilot processes. Characteristics of the primary effluent were quantified on samples from the inlet of the AGS-CFR. The average COD, TN, and TP were 340±90 mg/L, 55±5 mg/L, and 6.2±0.4 mg/L, respectively during the operation of configuration 1. The average COD, TN, and TP were 400±100 mg/L, 51±7 mg/L, and 6±1 mg/L, respectively during the operation of configuration 2.
Further characterization of the primary effluent was performed assuming that effluent filtered-flocculated COD was representative of the influent nbsCOD – Equation 15. This assumption was reasonable since the target and actual SRTs in both configurations were greater than 30 d [12]. Thus, the influent bsCOD could be estimated – Equation 16. If the influent nbpCOD was 0.08 for primary effluent, as typically assumed in conventional activated sludge models [13], the influent bpCOD could also be estimated – Equation 17. The estimations of nbsCOD and bsCOD were more reliable than for bpCOD, since bpCOD was not directly measured and depended on an assumption for nbpCOD. A summary of COD fractions is summarized in Table 9. Compared to typical fractions, the primary effluent supplied to the AGS-CFR had lower rapidly biodegradable COD, but in general the wastewater appeared to be more biodegradable overall (i.e., sum of bsCOD and bpCOD). Furthermore, the fraction of TN that was TAN was on average 0.72. The fraction of total phosphorus as OP was on average 0.71.

\[
\text{Equation 15} \\
nbsCOD = \frac{ff\text{COD}_{\text{effluent}}}{\text{COD}_{\text{influent}}}
\]

\[
\text{Equation 16} \\
bsCOD = \frac{ff\text{COD}_{\text{influent}} - ff\text{COD}_{\text{effluent}}}{\text{COD}_{\text{influent}}}
\]

\[
\text{Equation 17} \\
bpCOD = 1 - nbsCOD - bsCOD - 0.08
\]
Table 9 Measured/estimated characteristics of equalized primary effluent from a full-scale domestic wastewater treatment plant. Values shown for steady-state operation of configuration 1 and configuration 2 with one standard deviation. Measured/estimated values were compared to default values in conventional modelling software [13].

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Configuration 1</th>
<th>Configuration 2</th>
<th>BioWin default</th>
</tr>
</thead>
<tbody>
<tr>
<td>bsCOD</td>
<td>0.22 ± 0.04</td>
<td>0.20 ± 0.06</td>
<td>0.27</td>
</tr>
<tr>
<td>bpCOD</td>
<td>0.67 ± 0.05</td>
<td>0.66 ± 0.08</td>
<td>0.57</td>
</tr>
<tr>
<td>nbsCOD</td>
<td>0.04 ± 0.02</td>
<td>0.06 ± 0.03</td>
<td>0.08</td>
</tr>
<tr>
<td>nbpCOD</td>
<td>0.08 (assumed)</td>
<td>0.08 (assumed)</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Analytical methods and calculations

Influent and effluent samples were analyzed for COD, fCOD (i.e., after flocculation by zinc sulphate, adjustment to pH 10.5, 15 min sedimentation, and then 0.45 μm filtration; Nalgene syringe filter, Thermo Scientific, USA), TN, TP (TNTplus, Hach, CA), TAN, nitrite nitrogen, nitrate nitrogen, and OP (Lachat QuikChem 8500, HACH, CA). Samples for soluble nutrient analysis that required filtration were treated by medium porosity Q5 filter paper (Fisher Scientific, CA). MLSS, MLVSS, effluent TSS and VSS, and SVI were analyzed according to Standard Methods [14]. A SteREO Discovery (Zeiss, DE) was used for microscopic analysis. ImageJ software was used to assess the cross-sectional area and $D_F$ of particles with a cross-sectional area greater than or equal to 0.03 mm² (i.e., minimum diameter of 0.2 mm). $D_F$ is the ratio of particle perimeter and pi. The cross-sectional area was used to compute the diameter of a sphere with the same cross-sectional area (i.e., $D_s$). One-way analysis of variance (VassarStats, USA) was used to determine if steady-state data for configuration 1 was significantly different than for configuration 2 (i.e., $\alpha = 0.05$).
OLR was estimated based on Equation 1. It should be noted that the volume of the clarifier was neglected in OLR estimations. The actual SRT was estimated according to Equation 18. It should be noted that the mass of solids in the clarifier was neglected for SRT estimations, and therefore actual SRTs would be higher than estimated. The value for R was estimated according to Equation 19. Y_{net} was estimated according to Equation 20.

**Equation 18**

\[
SRT = \frac{MLSS_{aerobic} \cdot V_{anaerobic+aerobic} + MLSS_{anoxic} \cdot V_{anoxic}}{Q_{WAS} \cdot MLSS_{aerobic} + Q_{influent} \cdot TSS_{effluent}}
\]

**Equation 19**

\[
R = \frac{MLSS_{aerobic}}{MLSS_{anoxic} - MLSS_{aerobic}}
\]

**Equation 20**

\[
Y_{net} = \frac{Q_{WAS} \cdot MLSS_{aerobic} + Q_{influent} \cdot TSS_{effluent}}{Q_{influent} \cdot COD_{influent}}
\]

**Results**

**Reactor performance**

The AGS-CFR was inoculated in the summer (i.e., July 10, 2017) and data are presented for 270 d of operation. Temperature ranged from 16-25 °C and averaged 20±2 °C. The first 100 d were used to adjust operational and design aspects associated with DO control, hydraulics, overflows, aeration, and other parameters. As previously mentioned, ferric chloride was dosed to the full-scale primary clarifiers from 40-100 d. During ferric chloride dosing both the full-scale biological nutrient removal system and the AGS-CFR completely lost nitrification. This was apparent by the increase in effluent TAN from 40-100 d. After ferric chloride dosing was stopped, the AGS-CFR was setup in configuration 1 and re-seeded. Nitrification quickly
recovered, although on day 110 there was an aeration failure in the second aerobic tank that left
the second aerobic tank without air for 3 d. Consequently, nitrification was impacted and did not
fully recover until 140 d.

Interestingly, OP removal was not influenced by the aeration failure on day 110. In fact,
only nitrification was significantly impacted by the aeration failure. Instead, OP removal was
impacted by the accumulation of solids in the clarifier. During the beginning of the experiment
(i.e., 0-100 d), when the AGS-CFR was being tested for operational issues, the clarifier weirs
frequently clogged. The clarifier was successfully adjusted by day 40, which resulted in an
immediate drop in effluent OP concentrations. For the remaining 230 d of operation (i.e., 40-270
d), there were only a few instances where a scum layer accumulated in the clarifier and caused
increased effluent OP concentrations. Two of these instances occurred between 160-180 d and
resulted in approximately 0.5 mg-P/L effluent OP. Another two instances occurred between 200-
220 d and resulted in close to 1 mg-P/L effluent OP.

For configuration 1, steady-state was observed over 140-180 d. The aeration failure on
110 d delayed the onset of steady-state conditions for configuration 1. The time from 180-210 d
was not considered to contribute to steady-state for configuration 1 since the flow, and therefore
OLR, was increased to test process limits (i.e., not a part of the objectives for this paper). This
resulted in the HRT changing from 12 h to 6 h. However, nitrification was completely lost at the
6 h HRT and recovery of nitrification was not detected during the 30 d period at 6 h HRT. It was
then decided to stop operation in configuration 1, change to configuration 2, and re-seed. The
HRT was increased back to 12 h such that configuration 1 and 2 could be more accurately
compared. The steady-state period for configuration 2 was observed over 240-270 d.
Table 10 summarized effluent quality during steady-state operation for configuration 1 and configuration 2. In general, the performance of both configurations was good considering low-strength primary effluent was treated. Near complete removal of TAN was recorded and effluent OP averaged 0.2±0.2 mg-P/L in both configurations despite effluent TIN of 22±2 mg/L in configuration 1 and 17±2 mg/L in configuration 2. Configuration 1 produced higher effluent TSS and VSS, which were 23±20 mg/L and 20±20 mg/L respectively (i.e., VSS/TSS of 0.87), than configuration 2. Effluent TSS and VSS in configuration 2 averaged 16±9 mg/L and 14±8 mg/L, respectively (i.e., VSS/TSS of 0.88). Higher effluent VSS in configuration 1 negatively influenced effluent TP, which averaged 1.4±0.9 mg/L. Effluent TP in configuration 2 was only 0.7±0.2 mg/L. Based on effluent OP and TP concentrations, it was determined that effluent solids in configuration 1 contained 6% phosphorus (i.e., effluent particulate phosphorus divided by effluent VSS). Effluent solids in configuration 2 contained 3.6% phosphorus.

Table 10 Average steady-state effluent characteristics for configuration 1 and 2 shown with one standard deviation.

<table>
<thead>
<tr>
<th>Effluent (n = 5-17)</th>
<th>Configuration 1 (mg/L)</th>
<th>Configuration 2 (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSS</td>
<td>20 ± 20</td>
<td>14 ± 8</td>
</tr>
<tr>
<td>TSS</td>
<td>23 ± 20</td>
<td>16 ± 9</td>
</tr>
<tr>
<td>fCOD</td>
<td>14 ± 6</td>
<td>25 ± 10</td>
</tr>
<tr>
<td>COD</td>
<td>40 ± 20</td>
<td>50 ± 20</td>
</tr>
<tr>
<td>TAN</td>
<td>1 ± 0.6</td>
<td>0.5 ± 0.4</td>
</tr>
<tr>
<td>TIN</td>
<td>22 ± 2</td>
<td>17 ± 2</td>
</tr>
<tr>
<td>OP</td>
<td>0.2 ± 0.2</td>
<td>0.2 ± 0.2</td>
</tr>
<tr>
<td>TP</td>
<td>1.4 ± 0.9</td>
<td>0.7 ± 0.2</td>
</tr>
</tbody>
</table>
Effluent TSS and VSS were most likely higher in configuration 1 due to higher MLSS and MLVSS concentrations. The MLVSS in configuration 1 averaged 2.5±0.2 g/L while it averaged 1.7±0.3 g/L in configuration 2. This would have resulted in higher solids loading rate to the clarifier when configuration 1 was operated. Furthermore, the flow of primary effluent to configuration 1 was higher than to configuration 2 (i.e., configuration 1 had a larger total volume than configuration 2, but the same HRT of 12 h was maintained). The solids loading rate to the clarifier while configuration 1 was operated averaged 1 kg-TSS/(m²•d), while it averaged 0.5 kg-TSS/(m²•d) for configuration 2. It is interesting to note that the VSS/TSS of reactor biomass in configuration 1 was 0.81 and 0.77 in configuration 2. In both cases the reactor biomass had significantly lower VSS/TSS than effluent solids. Comparable results have been recorded in lab-scale reactors [3].

Due to similar general arrangement of configuration 1 and 2, which was to supply all primary effluent to a protected anaerobic zone (i.e., the pre-anoxic zone removed nitrites/nitrates via endogenous respiration), most of the influent carbon was utilized for enhanced biological phosphorus removal. Thus, TN removal depended completely upon endogenous denitrification in the pre-anoxic zone and simultaneous nitrification-denitrification in the aerobic zones. The COD/TN of primary effluent supplied to configuration 1 ranged from 5.3-8.0 (i.e., average of 6.4±0.8), while for configuration 2 it ranged from 4.4-11.5 (i.e., average of 8±2). Although the COD/TN between configurations did not vary significantly (i.e., P = 0.0572), the lower average COD/TN for configuration 1 may partially explain the 5 mg/L difference in effluent TIN between configuration 1 and 2. As well, it should be noted that influent TN averaged 54.5 mg/L for configuration 1 and 50.8 mg/L for configuration 2, which differed by 3.7 mg-N/L. However, the difference in primary effluent TN concentration supplied to configuration 1 and 2 was not
determined to be significantly different (i.e., \( P = 0.1067 \)). Thus, difference in average COD/TN and average TN of the primary influent may not have explained the statistically significant difference in effluent TIN between configurations (i.e., \( P < 0.0001 \)).

*Nitrogen and phosphorus profiles*

Steady-state concentrations of nitrogen and phosphorus species were monitored in each reactor zone – Figure 33. In general, trends were similar between configuration 1 and configuration 2. Primary effluent and RAS were pumped to the first anaerobic zone. A majority of the total phosphorus released occurred in the first anaerobic zone. The average OP concentration in the first anaerobic zone of configuration 1 was \( 31 \pm 6 \text{ mg-P/L} \) (i.e., 91% of total phosphorus release) and \( 32 \pm 3 \text{ mg-P/L} \) (i.e., 73% of total phosphorus release) in configuration 2. The OP concentration in the second anaerobic zone was \( 34 \pm 8 \text{ mg-P/L} \) in configuration 1 and \( 44 \pm 3 \text{ mg-P/L} \) in configuration 2. Nitrite and nitrate in both the first and second anaerobic zones was not detected in either configuration. As well, the concentration of TAN in both the first and second anaerobic zones was stable in either configuration, ranging from 20-22 mg/L on average.
Figure 33 Steady-state reactor nutrient profiles for configuration 1 and 2 without selective wasting. Error bars represent one standard deviation.
In the first aerobic zone, which was the biggest difference between configuration 1 and 2 (i.e., the first aerobic zone in configuration 1 was 140 L while it was only 30 L in configuration 2), there was a significant difference in nutrient profiles. For instance, in configuration 1 the TAN was 10±3 mg/L and nitrate was 9±6 mg-N/L, while nitrite was only 0.8±0.2 mg-N/L. The OP was also significantly removed in the first aerobic zone of configuration 1, averaging 2±3 mg-P/L. However, in configuration 2 there was significantly higher TAN in the first aerobic zone, which averaged 17±1 mg/L. Furthermore, there was less nitrate, at 0.6±0.5 mg-N/L, and more nitrite, at 2.6±0.4 mg-N/L, in the first aerobic zone of configuration 2. There was also less phosphorus uptake in the first aerobic zone of configuration 2, with OP averaging 24±4 mg-P/L. In general, these results indicated that the metabolic rates, and potentially metabolic pathways, in the first aerobic zone of configuration 1 and 2 were significantly different.

In the second aerobic zone and the final clarifier there was less of a difference between configuration 1 and configuration 2. The TAN in the second aerobic zone averaged 0.2±mg/L for configuration 1 and 0.2±0.2 mg/L for configuration 2. Similarly, TAN in the clarifier of configuration 1 averaged 0.3±0.4 mg/L while it averaged 0.3±0.2 mg/L for configuration 2. The average OP for both the second aerobic zone and clarifier, for both configurations, ranged from 0.1-0.2 mg-P/L and the maximum standard deviation was 0.2 mg-P/L. There was a significant difference in the nitrite, nitrate, and TIN concentrations between configurations 1 and 2. Nitrite averaged 0.4±0.2 mg-N/L and 0.5±0.1 mg-N/L in the second aerobic zone and clarifier, respectively, of configuration 1. In configuration 2 the nitrite was much higher, averaging 3.3±0.5 mg-N/L in the second aerobic zone and 3.4±0.5 mg-N/L in the clarifier. For this reason, the nitrate concentration in the second aerobic zone and clarifier of configuration 2. For instance,
nitrate was 20±3 mg-N/L in both the second aerobic zone and clarifier of configuration 1 while nitrate was 13±2 mg-N/L in both the second aerobic zone and clarifier of configuration 2.

Interestingly, there was a significant difference in TIN concentration (i.e., P <0.0001) for the second aerobic zone and clarifier between configuration 1 and 2. The TIN averaged 20±3 mg-N/L in the second aerobic zone and clarifier of configuration 1, while in the second aerobic zone and clarifier of configuration 2 TIN averaged 17±2 mg-N/L and 16±2 mg-N/L, respectively. The lower TIN in configuration 2 suggested that nitrogen removal was enhanced in configuration 2 compared to configuration 1. Furthermore, examination of the TIN profile for configuration 2 (i.e., Figure 1) clearly depicted a drop in TIN from the second anaerobic zone (i.e., 22±2 mg-N/L) to the clarifier (i.e., 16±2 mg-N/L). In configuration 1 there was no significant decrease in TIN from the second anaerobic zone (i.e., 20±1 mg-N/L) to the clarifier (i.e., 20±3 mg-N/L). Therefore, it can be concluded that configuration 2 had experienced simultaneous nitrification-denitrification while configuration 1 did not. As previously discussed, differences in average COD/TN and average TN of the primary influent supplied to configurations 1 and 2 were not significantly different, thus the significant drop of TIN across the aerobic zones in configuration 2 suggested that simultaneous nitrification-denitrification was the driving factor behind lower TIN in the effluent of configuration 2.

Lastly, there was also a significant difference between configuration 1 and 2 regarding nutrient concentrations in the pre-anoxic zone. Specifically, the difference was significant for nitrate and OP concentrations in the pre-anoxic zone. For example, nitrate in the pre-anoxic zone of configuration 1 averaged 9±3 mg-N/L, while it averaged below detection (i.e., detection limit of 0.2 mg-N/L) for configuration 2. The OP concentration in the pre-anoxic zone averaged 0.2±0.1 mg-P/L for configuration 1 and 3±3 mg-P/L for configuration 2. The inability of the pre-
anoxic zone in configuration 1 to fully remove nitrate versus the complete removal of nitrate in configuration 2 demonstrated that both residual carbon sources (e.g., polyhydroxyalkanoate) and the contact time for endogenous respiration were insufficient for full denitrification. Recalling that the pre-anoxic zone of configuration 1 was 100 L and configuration 2 was 120 L, the anoxic HRT of configuration 1 was 1.6 h while the anoxic HRT of configuration 2 was 2.2 h. On the other hand, the full removal of nitrate in the pre-anoxic zone of configuration 2 demonstrated that contact time and residual carbon sources were sufficient for full denitrification and fermentation, either to volatile fatty acid used by conventional PAO [15] or glucose used by species such as *Tetrasphaera* [16], which resulted in the release of OP.

**Biomass characteristics and morphology**

A summary of steady state operational parameters is displayed in Table 11. The MLSS in the second aerobic zone of configuration 1 stabilized at 3.1±0.3 g/L while in configuration 2 the MLSS stabilized at 2.2±0.4 g/L. Based on MLSS of the second aerobic zone alone, configuration 2 was 71% of the biomass in configuration 1. However, the MLSS of RAS, which was also the MLSS in the pre-anoxic zone, was 5.7±0.8 g/L in configuration 1 and 5.2±0.7 g/L in configuration 2. The difference in MLSS of RAS meant that the actual R was different between configurations. The target R was 1 in both cases, but the measured R was 1.3±0.2 in configuration 1 and 0.8±0.2 in configuration 2. Considering the respective volumes of individual zones, the overall mass of MLSS was 2.45±0.08 kg in configuration 1 and 1.9±0.1 kg in configuration 2. Thus, the biomass in configuration 2 was 78% of the biomass in configuration 1. The VSS/TSS in configuration 1 was 0.8±0.1, while it was 0.8±0.2 in configuration 2.
The 22% (i.e., 100% minus 78%) difference in weight of biomass between configuration 1 and 2 was not the result of changing the total volume between configuration 1 and configuration 2 because HRT of 12 h was targeted in both cases. Thus, the steady state OLR was similar between configuration 1, at 0.73±0.05 kg-COD/(m³•d), and configuration 2, at 0.8±0.2 kg-COD/(m³•d). Furthermore, there was no significant difference detected in the estimated SRT, which was 31 d in configuration 1 and 32 d in configuration 2. A reasonable explanation for the 22% difference in weight of biomass between configurations was that the ratio of anoxic, anaerobic, and aerobic volumes was significantly different between configurations, therefore resulting in different metabolisms. For instance, it is well known that anaerobic and anoxic yields are lower than aerobic yields [13]. Interestingly, Y\text{net} was estimated to be 0.15±0.08 kg-TSS/kg-COD in configuration 1 and 0.12±0.04 kg-TSS/kg-COD in configuration 2. Thus, Y\text{net} in configuration 2 was also less than in configuration 1 (i.e., 20% lower). However, the difference in Y\text{net} between configurations was not determined to be significantly different (i.e., P = 0.3695).
Steady state SVI in configuration 1 was 110±20 mL/g-MLSS (i.e., n = 13), while SVI in configuration 2 was 84±9 mL/g-MLSS (i.e., n = 11) – Figure 34. The difference in SVI between configurations was determined to be significantly different (i.e., P = 0.0006). Both configurations had significantly lower SVI than the control, which was 400±100 mL/g-MLSS (i.e., n = 3). Thus, both configuration 1 and configuration 2 significantly improved settleability of the inoculum, which was expected of an AGS process. Steady-state Ds of granular sludge in configuration 1 was 0.24±0.03 mm and the Df/Ds was 1.9±0.4 (i.e., n = 74). In configuration 2 the steady-state Ds of granular sludge was 0.3±0.1 mm and the Df/Ds was 1.8±0.3 (i.e., n = 109). Values for Df/Ds greater than 1 demonstrated that the granules in configuration 1 and 2 were slightly irregular.

The SVI of biomass in configuration 2 decreased significantly with the introduction of selective wasting (i.e., P <0.0001). Within 30 d the SVI decreased to <60 mL/g-MLSS, which is indicative of a well settling sludge. The maximum size of granules did not significantly change with selective wasting (i.e., max Ds in configuration 2 without selective wasting was 0.92 mm while it was 0.64 mm with selective wasting), but the presence of larger granules was much higher. Average Ds of granular sludge in configuration 2 with selective wasting, 12-48 d after the introduction of selective wasting, was 0.4±0.1 mm and the Df/Ds was 1.4±0.2. The lower Df/Ds value with selective wasting indicated that selective wasting resulted in more uniform and regularly shaped granules.
Figure 34 Top – steady-state SVI of configuration 1 and 2 without selective wasting (SW) and configuration 2 with selective wasting compared to inoculum. Error bars represented one standard deviation; Bottom – trend in SVI with the introduction of selective wasting to configuration 2.
Although the average Dₘ in configuration 1 and 2 seemed similar, they were determined to be significantly different (i.e., P = 0.0011). The difference in average Dₘ between configuration was due to the wider range of particle sizes in configuration 2 – Figure 35. For instance, the minimum, first quartile, median, third quartile, and maximum Dₘ in configuration 1 were 0.20 mm, 0.21 mm, 0.24 mm, 0.26 mm, and 0.36 mm, respectively. In configuration 2 the minimum, first quartile, median, third quartile, and maximum Dₘ were 0.20 mm, 0.23 mm, 0.25 mm, 0.29 mm, and 0.92 mm, respectively. Thus, 25% of granular sludge particles were between 0.26-0.36 mm in configuration 1, while 25% of granular sludge particles were between 0.29-0.92 mm in configuration 2.

Larger granules in configuration 2 may also be related to the simultaneous nitrification-denitrification observed in configuration 2. In fact, a recent study had suggested that the optimal range of Dₘ for nutrient removal by AGS was 0.28-0.45 mm [17]. In configuration 2, 29% of granular sludge particles fell within the optimal range while only 14% of the granular sludge particles in configuration 1 fell within the optimal range. Furthermore, considering that particles larger than 0.45 mm would also have contributed to simultaneous nitrification-denitrification, although not at the optimal rate [17], configuration 2 could have had 38% of the granular sludge particles contributing to simultaneous nitrification-denitrification.
The higher percentage of granules in the optimal range for simultaneous nitrification-denitrification may have explained why configuration 2 had better TIN removal resulting from a reduction of TIN across the aerobic zones. This would have also explained why configuration 2 with selective wasting had the best TN removal overall, since it had the highest proportion of large granules and therefore potentially the most simultaneous nitrification-denitrification - Figure 35. With selective wasting, the proportion of granules in the optimal range for simultaneous nitrification-denitrification was even higher. The minimum, first quartile, median, third quartile, and maximum Ds in configuration 2 with selective wasting were 0.22 mm, 0.26 mm, 0.28 mm, 0.42 mm, and 0.64 mm, respectively, 12-48 d after selective wasting was initiated. With selective wasting in configuration 2, 39% of the granules fell within the optimal range for simultaneous nitrification-denitrification while 56% of the granules could have been contributing to simultaneous nitrification-denitrification in general.
Figure 35 Top – particle size analysis for configuration 1, configuration 2 without selective wasting, and configuration 2 with selective wasting; Bottom – total nitrogen removal for configuration 1 (i.e., steady state), configuration 2 without selective wasting (i.e., steady state), and configuration 2 with selective wasting (i.e., 12-48 d after selective wasting was started). Error bars represented one standard deviation.
Summary

A mature aerobic granular sludge was cultivated within 40 d in a continuous flow reactor treating real primary effluent without VFA addition. Effluent OP was <0.2 mg/L and effluent TIN was <10 mg/L with 0.4±0.1 mm granules and a SVI <60 mL/g-MLVSS. Total nitrogen removal improved with the appearance and higher proportion of larger granules. Selective wasting was required to accumulate larger granules and wash-out the light, flocculent fraction of biomass. The pilot-scale AGS-CFR showed a high degree of performance and demonstrated that aerobic granular sludge is suitable for continuous flow applications involving low-strength, domestic wastewater following primary treatment. These results create the possibility for AGS technology to be applied to existing plug-flow type bioreactors.
Bibliography


Chapter 9: Modelling the AGS-CFR with BioWin™

Introduction

Although significant progress on AGS-CFRs for low-strength wastewater was made, there are still many future studies that could be performed. One of the most important would be to study the microbiome during and after granulation, as well as throughout seasonal operation. Analysis of the microbiome would be important to assess which microorganisms were present before and after granulation, and whether those microorganisms were quickly or gradually washed out, or if new microorganisms appeared rapidly or slowly. Information on the microbiome could be used to develop optimized start-up strategies that would result in efficient transition from flocculent to granular biomass. The information could also be used to re-evaluate existing activated sludge models such as BioWin™. The objective of this study was to see if the results from Chapter 8, without selective wasting, could be estimated by BioWin™. As well, parameters significantly impacting the modelled results were determined and described.

Methods

Results from Chapter 8 were first modelled with default BioWin™ kinetic and stoichiometric parameters (i.e., Appendix D). Default BioWin™ parameters were then modified to best match simulated results with the actual measured data from Chapter 6.
Results

Measured values from Chapter 8 were poorly modelled when default BioWin™ parameters were used – Table 12 and Figure 36. In fact, default BioWin™ parameters resulted in a predication that PAO would not be present in the AGS-CFR, although significant PAO activity was observed on the low-strength wastewater without VFA addition.

Table 12 Comparison of select parameters between actual measured data and an unoptimized and optimized BioWin™ model.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Effluent COD, mg/L</td>
<td>40</td>
<td>40</td>
<td>36</td>
</tr>
<tr>
<td>Effluent TIN, mg/L</td>
<td>22</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td>Effluent TAN, mg/L</td>
<td>1</td>
<td>0.1</td>
<td>0.4</td>
</tr>
<tr>
<td>Effluent TP, mg/L</td>
<td>1.4</td>
<td>4.9</td>
<td>1.7</td>
</tr>
<tr>
<td>Effluent OP, mg/L</td>
<td>0.2</td>
<td>4.5</td>
<td>0.0</td>
</tr>
<tr>
<td>MLSS, mg/L</td>
<td>3100</td>
<td>4510</td>
<td>4180</td>
</tr>
<tr>
<td>MLVSS, mg/L</td>
<td>2500</td>
<td>3890</td>
<td>2660</td>
</tr>
<tr>
<td>TSSRAS, mg/L</td>
<td>5700</td>
<td>7910</td>
<td>7490</td>
</tr>
<tr>
<td>VSSRAS, mg/L</td>
<td>4500</td>
<td>6470</td>
<td>4850</td>
</tr>
<tr>
<td>SRT, d</td>
<td>31</td>
<td>32.4</td>
<td>32.2</td>
</tr>
</tbody>
</table>
Iterative optimization uncovered multiple default BioWin™ parameters that did not represent the kinetics or stoichiometry of the AGS-CFR in Chapter 8. The set of optimized parameters clearly demonstrated that the differences between the actual measured data and BioWin™ default parameters converged on a few elements: 1) hydrolysis; 2) nitrifiers; 3) OHO; and 4) PAO – Table 13. For instance, the optimized parameters mostly regarded hydrolysis rates, fermentation kinetics, PAO kinetics, and the yields of OHO and PAO. It was expected that hydrolysis rates and fermentation kinetics were both changed, since hydrolysis and fermentation process both act to reduce complex COD to readily bioavailable compounds that are utilized by PAO.
Table 13 Summary of default and optimized BioWin™ parameters.

<table>
<thead>
<tr>
<th>Element</th>
<th>Parameter</th>
<th>BioWin default</th>
<th>Optimized value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrolysis</td>
<td>Hydrolysis rate, 1/d</td>
<td>2.1</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>Anoxic hydrolysis factor, -</td>
<td>0.28</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Anaerobic hydrolysis factor, -</td>
<td>0.04</td>
<td>3</td>
</tr>
<tr>
<td>Nitrifiers</td>
<td>AOB $\mu_{\text{max}}$, 1/d</td>
<td>0.9</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td>NOB $\mu_{\text{max}}$, 1/d</td>
<td>0.7</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td>Fermentation rate, 1/d</td>
<td>1.6</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Fermentation half sat., mg-COD/L</td>
<td>5</td>
<td>0.01</td>
</tr>
<tr>
<td></td>
<td>Aerobic yield, -</td>
<td>0.666</td>
<td>0.3</td>
</tr>
<tr>
<td>OHO</td>
<td>$\mu_{\text{max}}$, 1/d</td>
<td>0.95</td>
<td>1.9</td>
</tr>
<tr>
<td></td>
<td>Aerobic yield, -</td>
<td>0.639</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>VFA sequestration half sat., mg-COD/L</td>
<td>5</td>
<td>0.01</td>
</tr>
<tr>
<td>PAO</td>
<td>Aerobic P/PHA uptake, mg-P/mg-COD</td>
<td>0.93</td>
<td>1.05</td>
</tr>
<tr>
<td></td>
<td>Yield of PHA on sequestration, mg-P/mg-COD</td>
<td>0.889</td>
<td>0.99</td>
</tr>
</tbody>
</table>

Furthermore, it was not that surprising that both OHO and PAO parameters had to be changed, since more recent studies have alluded to a PAO that can ferment more complex COD itself [1]. For instance, members of the *Tetrasphaera* genus are known to ferment glucose before using it for glycogen storage. A single species performing the work of both fermentative OHO and PAO would explain why the yields of OHO and PAO had to be reduced by more than half of the default values, since a single species may perform the reaction in a single step (i.e., therefore approximately half the yield of OHO and PAO performing the same steps in series). BioWin™ does not account for a self-fermenting PAO, and thus OHO and PAO end up being of equal abundance in the optimized model (i.e., Figure 37), while instead it may only be one group of microorganisms. These results clearly demonstrated why metagenomic studies would be important to improve existing activated sludge models, such as by incorporating self-fermenting PAO.
Figure 37 Relative abundance of select microbial guilds from the optimized BioWin™ model.
Bibliography

Chapter 10: Engineering Significance

Overview of conclusions

Several conclusions were drawn from the research items contributing to this thesis. The following list is a brief overview of the major conclusions impacting engineering decisions:

1. Granulation occurred with low-strength substrate (i.e., COD <300 mg/L; COD:TP <50);
2. Hydrodynamic shear force, in excess of shear provided by aeration, was not required;
3. A minimum USAV of 0.2 cm/s per kg-COD/(m³•d) was recommended;
4. Wasting of granules stabilized AGS at high OLR and lower USAV;
5. Granulation was possible in CFRs with low-strength substrate;
6. F/M should not exceed 1.2 kg-COD/(kg-VSS•d) during granulation;
7. Selective wasting enhanced AGS structure, and frequency of granules, in a CFR; and
8. Properly balanced “feast” and “famine” conditions enhanced granulation in a CFR.

Engineering significance

Low-strength wastewater

With average influent of 340±30 mg/L COD, 42±5 mg/L TN, and 7±1 mg/L TP an AGS-SBR achieved effluent OP of 0.2±0.4 mg-P/L and effluent TIN of 17 mg-N/L. Aerobic granular sludge formed and matured, from conventional activated sludge inoculum, within 40 d while treating the synthetic proteinaceous medium. The AGS remained stable for the remaining 120 d experimentation period with an average F/M of 0.25±0.08 g-COD/(g-VSS•d). These results clearly demonstrated that properly designed AGS processes could overcome the barriers others have experienced with low-strength wastewater [1,2].
Low-strength wastewater was used in all studies besides that presented in Chapter 4, which examined high-strength brewery wastewater instead. All studies successfully cultivated AGS whether it was on synthetic proteinaceous medium or primary effluent from a full-scale domestic wastewater treatment facility. The full-scale primary effluent averaged 310±60 mg/L COD, 48±9 mg/L TN, and 5.7±0.9 mg/L TP and stable AGS did not lose performance despite COD as low as 269 mg/L and fCOD as low as 44 mg/L. Thus, in addition to the Chapter 2 confirming the suitability of AGS to treat low-strength wastewater, it was repeatedly demonstrated that granulation could be achieved on different sources of low-strength wastewater. These results are important because they provided repeatable instances of successful granulation on low-strength wastewater. Therefore, these results should provide confidence to those interested in using AGS to treat low-strength wastewaters, such as municipal wastewaters in North America. A summary of operational parameters for AGS treating low-strength wastewater is presented in Table 14.

Table 14 Summary of operational parameters for AGS treating low-strength wastewater.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLSS</td>
<td>4-12</td>
<td>g/L</td>
</tr>
<tr>
<td>HRT</td>
<td>6-12</td>
<td>h</td>
</tr>
<tr>
<td>OLR</td>
<td>0.5-1.5</td>
<td>kg-COD/(m³•d)</td>
</tr>
<tr>
<td>F/M</td>
<td>&lt;1.2</td>
<td>kg-COD/(kg-VSS•d)</td>
</tr>
<tr>
<td>SRT&lt;sub&gt;AGS&lt;/sub&gt;</td>
<td>30-50</td>
<td>d</td>
</tr>
<tr>
<td>SRT&lt;sub&gt;SW&lt;/sub&gt;</td>
<td>0.5-6</td>
<td>d</td>
</tr>
<tr>
<td>V&lt;sub&gt;critical&lt;/sub&gt;</td>
<td>2-8</td>
<td>cm/min</td>
</tr>
</tbody>
</table>
**Hydrodynamic shear force**

Chapter 3 examined the influence of increasing OLR on AGS-SBR operation under low hydrodynamic shear force. Specifically, USAV in the AGS-SBRs was 0.41 cm/s while OLR increased from 1.36 kg-COD/(m³•d) to 2.52 kg-COD/(m³•d) to 5.20 kg-COD/(m³•d). At 0.41 cm/s and 1.36 kg-COD/(m³•d), a mature AGS developed within 40 d. At OLR of 2.52 kg-COD/(m³•d) a mature AGS developed within a month, but filamentous microorganisms were detected on granule surfaces approximately 15 d after mature AGS development. Granules at OLR of 2.52 kg-COD/(m³•d) and USAV of 0.41 cm/s were fully coated by a filamentous surface layer within two months from inoculation. At OLR of 5.20 kg-COD/(m³•d) large granules developed rapidly, within two days, but then quickly destabilized. For the remaining four months of experimentation, biomass in the SBR with OLR of 5.20 kg-COD/(m³•d) and USAV of 0.41 cm/s appeared as filaments joined together by a white, viscous substance that was likely a mixture of EPS and some cells.

It was also demonstrated in Chapter 5 that wastewater treatment with nitrification would approach an A/A of 1 sooner than wastewater treatment without nitrification. This was because nitrification requires additional oxygen to biologically oxidize ammonium to nitrate. Therefore, USAV may not be an important parameter for AGS at municipal wastewater treatment facilities where complete nutrient removal was required. On the other hand, AGS may be less ideal for high-strength, organic wastewater with nitrogen deficiency (i.e., where nitrification would not be required). The reason being that additional aeration might be required to achieve the minimum USAV to guarantee AGS stability. However, it is possible that shear could be provided via different means, for example at high intensity for shorter duration, to minimize energy demand.
Solids retention time control at high organic loading rates

An alternative method for stabilizing AGS, compared to high hydrodynamic shear force, was examined in Chapter 4 while treating brewery wastewater. Specifically, the controlled wasting of AGS, in addition to the selective wasting of slowly-settling biomass, was examined at OLR from 1.5-4.5 kg-COD/(m³•d) and low USAV of 0.51 cm/s. The measured whole sludge SRT was approximately 7 d under all OLR. Previously, in Chapter 3, a minimum USAV of 0.2 cm/s per kg-COD/(m³•d) was recommended to maintain stable AGS. Thus, at 4.5 kg-COD/(m³•d) the USAV should have been 0.9 cm/s to generate stable AGS. However, it was demonstrated that controlled wasting of AGS helped maintain stable AGS under lower USAV.

Therefore, controlled wasting of AGS could allow for lower USAV, thereby reducing aeration requirements. For municipal wastewater treatment, where nitrification was required, the USAV could be achieved by the airflow required for wastewater treatment. However, additional aeration would be required for purely heterotrophic processes, such as AGS for brewery wastewater treatment. Thus, controlled wasting of AGS would be an effective strategy for wastewaters with high organic content and low nitrogen content, or for applications where nitrification was not required.
Selective wasting in continuous flow reactors

Chapter 7 examined the formation of AGS in a CFR treating low-strength, proteinaceous wastewater (i.e., COD = 370±30 mg/L; TN = 43±7 mg/L; and TP = 10±2 mg/L) under increasing selective pressure at OLR of 1.5 kg-COD/(m³•d). A continuously-stirred clarifier was used to out-select the light fraction of biomass at 20 °C by operating at Re between 1500-5000 (i.e., 7-25 rpm with a 5” propeller). Successful granulation was observed, although below 15 rpm a floculent-granular biomass was developed and above 20 rpm large granules with a filamentous population formed. A visual representation of the granulation process as a function of selective pressure (i.e., low = 7 rpm, high = 25 rpm) is presented in Figure 38.

Figure 38 Visual representation of granulation as a function of selective pressure in CFRs.
Results indicated that between 15-20 rpm, which corresponded with Re from 3000-4000, granulation would be optimized. At 15 rpm the F/M was 0.90±0.06 kg-COD/(kg-VSS•d), while at 20 rpm F/M was 14.1±0.8 kg-COD/(kg-VSS•d). This was due to higher biomass washout at 20 rpm, which resulted in MLSS of 150±20 mg/L while MLSS was 2100±200 mg/L at 15 rpm. It was suggested that a F/M of 1.2 kg-COD/(kg-VSS•d) not be exceeded during granulation – Table 14. Thus, the suggested F/M limit would have been induced between 15 and 20 rpm.

Although MLSS was low at 20 rpm, granules had good characteristics (i.e., smooth surface, low filamentous microorganisms) and De ranged from 0.21-0.67 mm. Values for De at 15 rpm only ranged from 0.21-0.23 mm. Thus, the optimal selective pressure that would have maintained enough MLSS for sufficient wastewater treatment, while also generating larger granules, was between 15-20 rpm.

These results were important because they demonstrated that granulation could be achieved in CFRs, while also demonstrating that the selective pressure must be carefully considered during start-up (i.e., conversion of flocculent biomass to granules). A difference of only 5 rpm, or 1000 regarding Re, resulted in significantly different biomass. Although granules larger than 0.2 mm are desired (i.e., diameter at 15 rpm), MLSS of 150 mg/L (i.e., MLSS at 20 rpm) is not. Thus, these results suggested that a stepwise approach to increasing selective pressure may be beneficial for start-up of full-scale AGS-CFR processes. By starting with low selective pressure, a hybrid flocculent-granular biomass would develop. The selective pressure could then be increased in increments such that the flocculent biomass was eventually washed out, resulting in a completely granular sludge system without a rapid decrease in MLSS. This is an important consideration for full-scale facilities since rapid loss of MLSS would likely correspond with poor performance and therefore low-quality effluent.
Feast and famine conditions in continuous flow reactors

The ratio of feast to famine conditions was studied in Chapter 8. An AGS-CFR was operated without selective wasting and two feast to famine ratios: 1) 0.47 v/v; and 2) 0.1 v/v. Primary effluent from a full-scale municipal wastewater treatment plant was supplied to the AGS-CFR. The OLR was 0.7-0.8 g-COD/(m$^3$ d). At lower feast to famine ratio, granulation was enhanced. For instance, SVI was 110±20 mL/g-VSS at feast to famine ratio of 0.47 while SVI was 86±8 mL/g-VSS at feast to famine ratio of 0.1. Both feast to famine ratios had significantly better SVI than the inoculum, which was 400±100 mL/g-VSS. This could be attributed to the overall reactor configuration, operated as anoxic/extended anaerobic/aerobic, versus the mainstream reactor, operated as anoxic/anaerobic/anoxic/aerobic with VFA supplement, where the inoculum originated. As well, granules were larger at lower feast to famine ratio. Values for D$s$ ranged from 0.20-0.36 mm at feast to famine ratio of 0.47, while D$s$ ranged from 0.20-0.92 mm at feast to famine ratio of 0.1. Therefore, it could be concluded that a short feast period followed by extended famine conditions benefited granulation in CFRs.

These results were important because they demonstrated design criteria for reactor configuration. Specifically, these results suggested that lower feast to famine ratios are beneficial for granulation, and therefore full-scale AGS-CFR facilities should be designed to ensure the development and transition from feast to famine conditions. In plug-flow reactors there will be a gradual transition, over reactor length, from feast to famine conditions. Defined barriers between feast and famine conditions in existing plug-flow reactors could be installed via baffles. Similarly, existing CMRs could be retrofitted by splitting existing basins into two or more basins, either by installing baffles or new tank walls.
**Summary of findings**

Experimental data clearly demonstrated that granulation, both in SBR and CFR, could be attained on low-strength wastewater. In fact, granulation was enhanced by extended famine conditions. Therefore low-strength wastewater may even improve granulation as long as adequate feast conditions develop. However, precautions should be taken to ensure that feast and famine conditions are properly balanced, thereby enhancing the granulation process. Feast and famine conditions represent a necessary, dynamic relationship between substrate concentrations and microorganisms required for successful and stable granulation. During feast conditions, organic substrate concentrations are high, representing a surplus of available energy compared to what would be required for biomass growth. As a result, the system, under feast conditions, favours the growth of EPS producing microbial strains, which invest a significant portion of the energy they capture into EPS production. The EPS may help initiate and mature microbial adhesion, whether through EPS mediated tethering or by increasing the attractive force between cells, AGS aggregates, and/or other organic material that may otherwise be hydrophilic. Therefore, feast conditions represent a necessary step towards the development of strong and stable biofilms, including AGS.

Extended famine conditions are also important and are required to balance the EPS production phase during feast conditions. During famine conditions, organic substrate concentrations are low, representing energy-limited conditions and therefore low availability of energy for growth. During famine conditions, microorganisms may consume some fractions of the EPS produced during the feast phase. As a result, the EPS matrix is modified, potentially increasing porosity of the EPS matrix which subsequently enhances diffusion rates. Adequate diffusion is required for anaerobic substrate utilization, thus extended famine conditions may,
indirectly, improve anaerobic substrate utilization. Extended famine conditions may also help slow-growing microorganisms out-compete fast-growing microorganisms through endogenous respiration mediated selective pressures. During extended famine conditions, microorganisms must rely on internal storage product to perform basic cellular maintenance functions. Fast-growing microorganisms, which are typically associated with higher decay rates, would be negatively impacted during extended famine conditions and their viable population would be significantly reduced. Slow-growing microorganisms, especially slow-growing microorganism capable of maintaining enough internal storage product reserves such as PAO, would be able to keep basic cellular maintenance functions active during extended famine conditions. Therefore, their population would not be significantly impacted by extended famine conditions. Extended famine conditions are a necessary step towards stable AGS, both structurally and for maintaining a desirable microbial community.

It was demonstrated that selective pressure also plays a role in the success of granulation for CFRs. During the conversion of existing, flocculent CFR processes to AGS, the selective pressure should be gradually increased, thereby maintaining high concentrations of active biomass leading to high effluent quality. This would allow for the gradual development of granules, ultimately resulting in AGS replacing flocculent biomass. The selective wasting process could be any physical process capable of separating AGS from flocculent biomass, as well as other low-density, undesirable matter. For instance, selective wasting can be induced by differential settling, such as in gravity settlers or hydrocyclones, or by size exclusion, such as sieves. It would be desirable for the selective wasting process be as flexible as possible, allowing for different selective pressures and wasting regimes to be employed.
For this reason, size exclusion based on sieve size may be the least appealing. Similarly, hydrocyclones would have a narrow range of operating conditions. Therefore, gravity settling in batch mode may be the most attractive. Batch methods allow for flexibility by changing the number of wasting cycles per day, as well as the volume wasted per cycle. Batch gravity settling would also provide flexibility in terms of the selective pressure applied. For instance, the time allocated for settling could be adjusted based on real-time process parameters. If a higher proportion of granules was desired, the settling time could be shortened and/or the frequency of wasting cycles increased, thereby increasing the selective pressure applied. On the other hand, if excessive washout was observed, the settling time could be increased and/or the frequency of wasting cycles decreased. The well-engineered use of selective wasting, coupled with adequately balanced feast and famine conditions, has great potential for integrating AGS into CFRs.

Lastly, it was demonstrated that high hydrodynamic shear force was not a requirement for granulation, and that the airflow provided for nitrification facilities would be enough to maintain AGS stability based on USAV. If additional attrition was required to stabilize AGS, it could be a process, aeration or otherwise, that provided shear at high intensity for a shorter duration immediately before the selective wasting process. As long as the shear was of sufficient time-intensity to provide the necessary attrition for scouring of undesirable matter from the surface of AGS. Thereby, AGS particles would be scoured of undesirable surface coatings immediately before selective wasting, allowing for the undesirable materials, whether particulate organic matter or filamentous microorganisms, to be removed with the selective waste. Furthermore, it was demonstrated that lower airflow, and therefore USAV, could be used if controlled wasting of AGS was employed. This could represent significant savings in electricity requirements for aeration at industrial facilities.
Recommendations for future studies

Although significant progress on AGS-CFRs for low-strength wastewater was made, there are still many future studies that can be performed. The most pertinent are discussed below:

1. Study microbiome during and after granulation

   It would be important to study the microbiome during and after granulation to assess which microorganisms were present before and after granulation, and whether those microorganisms were quickly or gradually washed out, or if new microorganisms appeared rapidly or slowly. That information could then be used to develop optimized start-up strategies that would result in efficient transition from flocculent to granular biomass. The information could also be used to improve existing activated sludge models. Chapter 9 clearly demonstrated that default kinetic and stoichiometric parameters in BioWin™ did not adequately represent the AGS-CFR from Chapter 8.

2. Study EPS fractions and incorporate into activated sludge models

   Another critical factor to the formation and stability of AGS is the development and maintenance of an EPS matrix. As of now, many mainstream activated sludge models, including BioWin™, do not consider the formation and breakdown of EPS components. However, some do consider the formation of storage compounds and the generation of microbial product from endogenous respiration – Figure 39. The examination of EPS production and modification in AGS could lead to modelled parameters that track the generation of AGS, and even biofilm, processes. Furthermore, the integration of EPS production and modification mechanisms into activated sludge models could result in more accurate predictions of AGS particle size, as well as biofilm thickness.
3. Develop and optimize a control system

Lastly, it was demonstrated that successful implementation of an AGS-CFR would require the development and maintenance of defined feast and famine conditions. Thus, a control system should be developed that can track feast and famine conditions in real-time, as well as relay online measurements to operational parameters such as the variable frequency drive on blowers, to standby blowers of swing zones, to the RAS pump, and other parameters. Stable granulation depends on the constant formation and maintenance of new and old AGS, thus control systems that monitor system performance in these regards are critical to both SBR and CFR applications of AGS.
Bibliography


Appendix A

Determination of $D_F$ and $D_S$

ImageJ is a free program package for image processing that can be downloaded from http://rsb.info.nih.gov/ij/download.html. ImageJ can be used to assess particle size distribution. The following steps were taken to obtain reliable results:

1. Set scale – A scale must be provided to obtain measurements in units other than pixels.
   Go to **Analyze > Set Scale**. Input the known distance, which is used to convert between pixels and the desired unit. The known distance can be determined from a reference item, such as the scale bar on microscopic images.

2. Remove the background – In most cases the background can interfere with ImageJ’s capability to differentiate between discrete particles and noise. In the case of granular sludge, the particles are granules and background noise can originate from flocculent or filamentous biomass. The noise can be reduced by removing the background. Go to **Image > Duplicate**. On the duplicated image go to **Process > Filters > Gaussian Blurr** and apply a value for radius between 20-50 pixels (i.e., value may change depending on substrate, lighting conditions, etc.). The Gaussian blurr converts the duplicate image to an image of just the background, which can then be removed from the original image. Go to **Process > Image Calculator** and subtract the duplicated image from the original image. The resulting image should clearly show the granules with reduced background noise.

3. Adjust the brightness – Go to **Image > Adjust > Brightness/Contrast** and select automatic. Manually adjust the brightness afterwards such that the brightness/contrast between granules and the background is optimized.
4. Convert to 8-bit – Go to **Image > Type > 8-bit.** This converts the image to greyscale, which is required for the next step.

5. Find a threshold for the image – Go to **Image > Adjust > Threshold.** The sliders must be adjusted in such a way that the granules appear as dark spots in the black and white mode.

6. Remove scale, fill in particles, remove “bad” particles – The image should now appear completely in black and white. Therefore, the paintbrush and paint-can icons can be used to remove the scale bar, fill in particles, and remove “bad” particles. To remove the scale bar, convert it to white. Particles can be filled in black with the paintbrush or paint-can. “Bad” particles, which mostly refers to particles existing on the border (i.e., total area cannot be measured, and their inclusion will skew results) can be removed by converting them to white.

7. Reduce surface irregularities and make sure particles are discrete – Go to **Process > Binary > Erode.** The erode function can be used to remove surface irregularities, such as filamentous microorganisms. However, use of the erode function may also require some back filling of the particles (i.e., paintbrush or paint-can black). Lastly, overlapping granules should be adjusted such that each particle is discrete, and therefore can be measured as a single particle. Use the white paintbrush with a thin line to separate particles.

8. Analyze the particles – Go to **Analyze > Set Measurements** and select Area and Feret’s diameter. Go to **Analyze > Analyze Particles.** Set the lower limit of Size to the cutoff diameter (i.e., if the cutoff diameter is 0.2 mm, the cutoff size is the corresponding area of 0.0314 mm²) and select ok.
A sample analysis for granular sludge, according to the described steps, is presented below:

Step 1.
Scale bar of 1 mm was determined to be 150 pixels.

Step 2.

minus equals
Step 3.

Step 4.
Appendix B

One-way ANOVA

One-way analysis of variance (ANOVA) tests the null hypothesis (i.e., $H_0 =$ there is no relationship between two or more measured phenomena) by analyzing the differences among group means in a sample. It assumes independence, normality, and homoscedasticity.

“VassarStats: Website for Statistical Computation” was used to perform one-way ANOVA (i.e., http://www.vassarstats.net). Sample calculations are provided for a scenario where three parallel bioreactors, which were identical besides the pH (i.e., pH controlled at 7, 8, and 9), were treating the same wastewater and effluent nitrite was quantified (i.e., $H_0 =$ there is no relationship between pH and effluent nitrite). Steady-state data is presented below:

<table>
<thead>
<tr>
<th>Day of steady-state</th>
<th>pH = 7</th>
<th>pH = 8</th>
<th>pH = 9</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.2</td>
<td>0.6</td>
<td>4.3</td>
</tr>
<tr>
<td>3</td>
<td>0.2</td>
<td>0.5</td>
<td>5.2</td>
</tr>
<tr>
<td>5</td>
<td>0.3</td>
<td>0.6</td>
<td>4.8</td>
</tr>
<tr>
<td>8</td>
<td>0.2</td>
<td>0.4</td>
<td>4.9</td>
</tr>
<tr>
<td>10</td>
<td>0.4</td>
<td>0.7</td>
<td>5.0</td>
</tr>
<tr>
<td>12</td>
<td>0.2</td>
<td>0.5</td>
<td>5.2</td>
</tr>
<tr>
<td>15</td>
<td>0.2</td>
<td>0.6</td>
<td>4.7</td>
</tr>
</tbody>
</table>

$P < 0.0001$ (i.e., $\alpha = 0.05$), therefore $H_0$ was rejected and there is a significant relationship between reactor pH and effluent nitrite. A print-out of the analysis is presented on the following pages.
One-Way Analysis of Variance for Independent or Correlated Samples

[Traducción en español] The logic and computational details of the one-way ANOVA for independent and correlated samples are described in Chapters 13, 14, and 15 of Concepts and Applications.

Procedure:

- **Initial Setup:**
  Enter the number of samples in your analysis (2, 3, 4, or 5) into the designated text field, then click the «Setup» button for either Independent Samples or Correlated Samples to indicate which version of the one-way ANOVA you wish to perform.

- **Entering Data Directly into the Text Fields:**
  After clicking the cursor into the scrollable text area for Sample 1, enter the values for that sample in sequence, pressing the carriage return key after each entry except the last. (On a Macintosh platform, the carriage return key is labeled 'Return'; on a Windows platform it is labeled 'Enter'.) Perform the same procedure for the other samples in your analysis.

- **Importing Data via Copy & Paste:**
  Within the spreadsheet application or other source of your data, select and copy the column of data for sample 1. Then return to your web browser, click the cursor into the text area for sample 1 and perform the 'Paste' operation from the 'Edit' menu. Perform the same procedure for the other samples in your analysis.

- **Data Check:**
  For each sample, make sure that the final entry is **not** followed by a carriage return. (A carriage return after the final entry in a sample will be interpreted as an extra data entry whose value is zero. Importing data via the copy and paste procedure will almost always produce an extra carriage return at the end of a column.) After all values for a sample have been entered, click the cursor immediately to the right of the final entry in the list, then press the down-arrow key. If an extra line is present, the cursor will move downward. Extra lines can be removed by pressing the down arrow key until the cursor no longer moves, and then pressing the 'Backspace' key (on a Mac platform, 'delete') until the cursor stands immediately to the right of the final entry.

  If you are performing a correlated-samples analysis, also make sure that the values for each sample are entered in the appropriate sequence. Note that a correlated-samples analysis presupposes equal numbers of observations for each sample in the analysis.

- **When all** sample values have been entered, click the button labeled «Calculate.» For independent samples the default analysis is a standard weighted-means analysis. If you wish to perform an unweighted-means analysis, click the «Unweighted» button before calculating.

Note that when the number of samples is \( k = 2 \), the analysis of variance (standard weighted-means analysis) is equivalent to a non-directional \( t \)-test with \( F = t^2 \).
### Data Entry

<table>
<thead>
<tr>
<th>Sample 1</th>
<th>Sample 2</th>
<th>Sample 3</th>
<th>Sample 4</th>
<th>Sample 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.2</td>
<td>0.6</td>
<td>4.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0.5</td>
<td>5.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3</td>
<td>0.6</td>
<td>4.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0.4</td>
<td>4.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.4</td>
<td>0.7</td>
<td>5.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0.5</td>
<td>5.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2</td>
<td>0.6</td>
<td>4.7</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Data Summary

<table>
<thead>
<tr>
<th></th>
<th>Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>N</td>
<td>7</td>
</tr>
<tr>
<td>-ΣX</td>
<td>1.7</td>
</tr>
<tr>
<td>Mean</td>
<td>0.242857</td>
</tr>
<tr>
<td>-ΣX²</td>
<td>0.45</td>
</tr>
<tr>
<td>Variance</td>
<td>0.00619</td>
</tr>
<tr>
<td>Std.Dev.</td>
<td>0.07668</td>
</tr>
<tr>
<td>Std.Err.</td>
<td>0.029738</td>
</tr>
</tbody>
</table>

ANOVA Summary: Independent Samples k=3

<table>
<thead>
<tr>
<th>Source</th>
<th>SS</th>
<th>df</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment [between groups]</td>
<td>93.649524</td>
<td>2</td>
<td>46.824762</td>
<td>1224.05</td>
<td>&lt;.0001</td>
</tr>
</tbody>
</table>
8/29/2018  

One-Way ANOVA

<table>
<thead>
<tr>
<th>Error</th>
<th>0.668571</th>
<th>18</th>
<th>0.038254</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ss/Bl</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>94.338095</td>
<td>20</td>
<td></td>
</tr>
</tbody>
</table>

Ss/Bl = Subjects or Blocks depending on the design.
Applicable only to correlated-samples ANOVA.

---

**Tukey HSD Test**

<table>
<thead>
<tr>
<th></th>
<th>HSD[.05] = 0.27; HSD[.01] = 0.35</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1 vs M2</td>
<td>P &lt; .05</td>
</tr>
<tr>
<td>M1 vs M3</td>
<td>P &lt; .01</td>
</tr>
<tr>
<td>M2 vs M3</td>
<td>P &lt; .01</td>
</tr>
</tbody>
</table>

M1 = mean of Sample 1
M2 = mean of Sample 2
and so forth.

HSD = the absolute [unsigned] difference between any two sample means required for significance at the designated level. HSD[.05] for the .05 level; HSD[.01] for the .01 level.

---

*[Link to page](http://www.vassarstats.net/)*

---

© Richard Lowry 2001-2018
All rights reserved.
Appendix C

Determination of $r_{\text{max}}$

Values for $r_{\text{max}}$ were determined from the maximum slope of concentration for the respective nutrient (i.e., orthophosphate or nitrite/nitrate) and mixed liquor volatile suspended solids in the reactor during kinetic tests, according to the following equation:

$$r_{\text{max}} = \frac{m_{\text{max, linear regression}}}{MLVSS_{\text{reactor}}}$$

The following discussion will provide examples for the determination of $r_{\text{max, NOx-N prod.}}$. During start-up (i.e., first month), NOx-N was produced at a constant rate throughout aeration:

Therefore, the maximum slope was 0.1337 mg-N/L per min. Converting to hours, the slope became approximately 8 mg-N/L per h. At that point in time, the mixed liquor volatile suspended solids were 2.3 g/L. Therefore, $r_{\text{max, NOx-N prod.}}$ was equal to 8 divided by 2.3, or approximately 3.5 mg-N/(g-VSS•h).
As the reactors began to approach steady-state, the NO\textsubscript{x}-N concentration would plateau about 60 min into aeration:

![Graph showing NO\textsubscript{x}-N concentration vs. aerobic time]

\[y = 0.2143x + 3.7733\]

\[R^2 = 0.9939\]

Thus, the maximum slope was calculated by linear regression only for points contributing to the maximum recorded slope. In this example, the maximum slope was 0.2143 mg-N/L per min, or approximately 13 mg-N/L per h. At that point in time, the mixed liquor volatile suspended solids were 4.5 g/L. Therefore, \( \Gamma_{\text{max, NOx-N prod.}} \) was equal to 13 divided by 4.5, or approximately 3 mg-N/(g-VSS•h).
Appendix D

BioWin™ model of the AGS-CFR

The AGS-CFR configuration was first set-up in BioWin™:

Next, the appropriate reactor volumes were incorporated, along with operational parameters (e.g., dissolved oxygen values, recycle ratios, waste activated sludge flows). The influent element was adjusted to steady-state concentrations and fractions for the full-scale primary effluent. Steady-state simulations were run, and a single kinetic or stoichiometric parameter was changed until the optimal output (i.e., closest matching to measured data) was obtained. A report of the optimized BioWin™ is provided on the following pages.
BioWin user and configuration data

Project details

Project name: AGS-CFR simulation       Project ref.: PhD AF
Plant name: UoM                        User name: TRD

Created: 17/08/2018                    Saved: 29/08/2018

Steady state solution

Total SRT: 32.25 days
Temperature: 20.0°C

Flowsheet
## Album page – Reactor profile

<table>
<thead>
<tr>
<th></th>
<th>Pre-anoxic</th>
<th>Anaerobic 1</th>
<th>Anaerobic 2</th>
<th>Aerobic 1</th>
<th>Aerobic 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ammonia N</td>
<td>28</td>
<td>23</td>
<td>22</td>
<td>21</td>
<td>21</td>
</tr>
<tr>
<td>Nitrite N</td>
<td>19</td>
<td>18</td>
<td>17</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Nitrate N</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Total inorganic N</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Soluble PO4-P</td>
<td>0.7</td>
<td>0.8</td>
<td>0.9</td>
<td>1.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>

*Note: Concentrations are in mg/L.*
Global Parameters

**Common**

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrolysis rate [1/d]</td>
<td>2.1000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Hydrolysis half sat. [-]</td>
<td>0.0600</td>
<td>0.0600</td>
</tr>
<tr>
<td>Anoxic hydrolysis factor [-]</td>
<td>0.2800</td>
<td>3.0000</td>
</tr>
<tr>
<td>Anaerobic hydrolysis factor (AS) [-]</td>
<td>0.0400</td>
<td>3.0000</td>
</tr>
<tr>
<td>Anaerobic hydrolysis factor (AD) [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Adsorption rate of colloids [L/(mgCOD d)]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
<tr>
<td>Ammonification rate [L/(mgCOD d)]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>Assimilative nitrate/nitrite reduction rate [1/d]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Endogenous products decay rate [1/d]</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
### AOB

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>0.9000</td>
<td>0.6000</td>
</tr>
<tr>
<td>Substrate (NH4) half sat. [mgN/L]</td>
<td>0.7000</td>
<td>0.7000</td>
</tr>
<tr>
<td>Byproduct NH4 logistic slope [-]</td>
<td>50.0000</td>
<td>50.0000</td>
</tr>
<tr>
<td>Byproduct NH4 inflection point [mgN/L]</td>
<td>1.4000</td>
<td>1.4000</td>
</tr>
<tr>
<td>AOB denite DO half sat. [mg/L]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>AOB denite HNO2 half sat. [mgN/L]</td>
<td>5.000E-6</td>
<td>5.000E-6</td>
</tr>
<tr>
<td>Aerobic decay rate [1/d]</td>
<td>0.1700</td>
<td>0.1700</td>
</tr>
<tr>
<td>Anoxic/anaerobic decay rate [1/d]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>KiHNO2 [mmol/L]</td>
<td>0.0050</td>
<td>0.0050</td>
</tr>
</tbody>
</table>

### NOB

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>0.7000</td>
<td>0.6000</td>
</tr>
<tr>
<td>Substrate (NO2) half sat. [mgN/L]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Aerobic decay rate [1/d]</td>
<td>0.1700</td>
<td>0.1700</td>
</tr>
<tr>
<td>Anoxic/anaerobic decay rate [1/d]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>KiNH3 [mmol/L]</td>
<td>0.0750</td>
<td>0.0750</td>
</tr>
</tbody>
</table>

### AAO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>0.2000</td>
<td>0.2000</td>
</tr>
<tr>
<td>Substrate (NH4) half sat. [mgN/L]</td>
<td>2.0000</td>
<td>2.0000</td>
</tr>
<tr>
<td>Substrate (NO2) half sat. [mgN/L]</td>
<td>1.0000</td>
<td>1.0000</td>
</tr>
<tr>
<td>Aerobic decay rate [1/d]</td>
<td>0.0190</td>
<td>0.0190</td>
</tr>
<tr>
<td>Anoxic/anaerobic decay rate [1/d]</td>
<td>0.0095</td>
<td>0.0095</td>
</tr>
<tr>
<td>Ki Nitrite [mgN/L]</td>
<td>1000.000</td>
<td>1000.000</td>
</tr>
<tr>
<td>Nitrite sensitivity constant [L / (d mgN)]</td>
<td>0.0160</td>
<td>0.0160</td>
</tr>
</tbody>
</table>
### OHO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate half sat. [mgCOD/L]</td>
<td>5.0000</td>
<td>5.0000</td>
</tr>
<tr>
<td>Anoxic growth factor [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Denite N2 producers (NO3 or NO2) [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Aerobic decay rate [1/d]</td>
<td>0.6200</td>
<td>0.6200</td>
</tr>
<tr>
<td>Anoxic decay rate [1/d]</td>
<td>0.2330</td>
<td>0.2330</td>
</tr>
<tr>
<td>Anaerobic decay rate [1/d]</td>
<td>0.1310</td>
<td>0.1310</td>
</tr>
<tr>
<td>Fermentation rate [1/d]</td>
<td>1.6000</td>
<td>3.0000</td>
</tr>
<tr>
<td>Fermentation half sat. [mgCOD/L]</td>
<td>5.0000</td>
<td>0.0100</td>
</tr>
<tr>
<td>Fermentation growth factor (AS) [-]</td>
<td>0.2500</td>
<td>0.2500</td>
</tr>
<tr>
<td>Free nitrous acid inhibition [mol/L]</td>
<td>1.000E-7</td>
<td>1.000E-7</td>
</tr>
</tbody>
</table>

### Methylotrophs

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>1.3000</td>
<td>1.3000</td>
</tr>
<tr>
<td>Methanol half sat. [mgCOD/L]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Denite N2 producers (NO3 or NO2) [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Aerobic decay rate [1/d]</td>
<td>0.0400</td>
<td>0.0400</td>
</tr>
<tr>
<td>Anoxic/anaerobic decay rate [1/d]</td>
<td>0.0300</td>
<td>0.0300</td>
</tr>
<tr>
<td>Free nitrous acid inhibition [mmol/L]</td>
<td>1.000E-7</td>
<td>1.000E-7</td>
</tr>
</tbody>
</table>

### PAO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>0.9500</td>
<td>1.9000</td>
</tr>
<tr>
<td>Max. spec. growth rate, P-limited [1/d]</td>
<td>0.4200</td>
<td>0.4200</td>
</tr>
<tr>
<td>Substrate half sat. [mgCOD(PHB)/mgCOD(Zbp)]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Substrate half sat., P-limited [mgCOD(PHB)/mgCOD(Zbp)]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>Acetogens</td>
<td>Default</td>
<td>Value</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>---------</td>
<td>-------</td>
</tr>
<tr>
<td>Max. spec. growth rate [1/d]</td>
<td>0.2500</td>
<td>0.2500 1.0290</td>
</tr>
<tr>
<td>Substrate half sat. [mgCOD/L]</td>
<td>10.0000</td>
<td>10.0000 1.0000</td>
</tr>
<tr>
<td>Acetate inhibition [mgCOD/L]</td>
<td>10000.0000</td>
<td>10000.0000 1.0000</td>
</tr>
<tr>
<td>Anaerobic decay rate [1/d]</td>
<td>0.0500</td>
<td>0.0500 1.0290</td>
</tr>
<tr>
<td>Aerobic/anoxic decay rate [1/d]</td>
<td>0.5200</td>
<td>0.5200 1.0290</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Methanogens</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetoclastic max. spec. growth rate [1/d]</td>
<td>0.3000</td>
<td>0.3000 1.0290</td>
</tr>
<tr>
<td>H2-utilizing max. spec. growth rate [1/d]</td>
<td>1.4000</td>
<td>1.4000 1.0290</td>
</tr>
<tr>
<td>Acetoclastic substrate half sat. [mgCOD/L]</td>
<td>100.0000</td>
<td>100.0000 1.0000</td>
</tr>
<tr>
<td>Acetoclastic methanol half sat. [mgCOD/L]</td>
<td>0.5000</td>
<td>0.5000 1.0000</td>
</tr>
<tr>
<td>H2-utilizing CO2 half sat. [mmol/L]</td>
<td>0.1000</td>
<td>0.1000 1.0000</td>
</tr>
<tr>
<td>H2-utilizing substrate half sat. [mgCOD/L]</td>
<td>1.0000</td>
<td>1.0000 1.0000</td>
</tr>
<tr>
<td>H2-utilizing methanol half sat. [mgCOD/L]</td>
<td>0.5000</td>
<td>0.5000 1.0000</td>
</tr>
<tr>
<td>Acetoclastic propionic inhibition [mgCOD/L]</td>
<td>10000.0000</td>
<td>10000.0000 1.0000</td>
</tr>
<tr>
<td>Acetoclastic anaerobic decay rate [1/d]</td>
<td>0.1300</td>
<td>0.1300 1.0290</td>
</tr>
<tr>
<td>Name</td>
<td>Default</td>
<td>Value</td>
</tr>
<tr>
<td>----------------------------------------------------------------------</td>
<td>---------</td>
<td>-------</td>
</tr>
<tr>
<td>Acetoclastic aerobic/anoxic decay rate</td>
<td>0.6000</td>
<td>0.6000</td>
</tr>
<tr>
<td>H2-utilizing anaerobic decay rate</td>
<td>0.1300</td>
<td>0.1300</td>
</tr>
<tr>
<td>H2-utilizing aerobic/anoxic decay rate</td>
<td>2.8000</td>
<td>2.8000</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>pH Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>OHO low pH limit [-]</td>
<td>4.0000</td>
<td>4.0000</td>
</tr>
<tr>
<td>OHO high pH limit [-]</td>
<td>10.0000</td>
<td>10.0000</td>
</tr>
<tr>
<td>Methylotrophs low pH limit [-]</td>
<td>4.0000</td>
<td>4.0000</td>
</tr>
<tr>
<td>Methylotrophs high pH limit [-]</td>
<td>10.0000</td>
<td>10.0000</td>
</tr>
<tr>
<td>Autotrophs low pH limit [-]</td>
<td>5.5000</td>
<td>5.5000</td>
</tr>
<tr>
<td>Autotrophs high pH limit [-]</td>
<td>9.5000</td>
<td>9.5000</td>
</tr>
<tr>
<td>PAO low pH limit [-]</td>
<td>4.0000</td>
<td>4.0000</td>
</tr>
<tr>
<td>PAO high pH limit [-]</td>
<td>10.0000</td>
<td>10.0000</td>
</tr>
<tr>
<td>OHO low pH limit (anaerobic)</td>
<td>5.5000</td>
<td>5.5000</td>
</tr>
<tr>
<td>OHO high pH limit (anaerobic)</td>
<td>8.5000</td>
<td>8.5000</td>
</tr>
<tr>
<td>Propionic acetogens low pH limit [-]</td>
<td>4.0000</td>
<td>4.0000</td>
</tr>
<tr>
<td>Propionic acetogens high pH limit [-]</td>
<td>10.0000</td>
<td>10.0000</td>
</tr>
<tr>
<td>Acetoclastic methanogens low pH limit [-]</td>
<td>5.0000</td>
<td>5.0000</td>
</tr>
<tr>
<td>Acetoclastic methanogens high pH limit [-]</td>
<td>9.0000</td>
<td>9.0000</td>
</tr>
<tr>
<td>H2-utilizing methanogens low pH limit [-]</td>
<td>5.0000</td>
<td>5.0000</td>
</tr>
<tr>
<td>H2-utilizing methanogens high pH limit [-]</td>
<td>9.0000</td>
<td>9.0000</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Switches Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>OHO DO half sat. [mgO2/L]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>PAO DO half sat. [mgO2/L]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>Anoxic/anaerobic NOx half sat. [mgN/L]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
<tr>
<td>AOB DO half sat. [mgO2/L]</td>
<td>0.2500</td>
<td>0.2500</td>
</tr>
<tr>
<td>Parameter</td>
<td>Default Value</td>
<td>Value</td>
</tr>
<tr>
<td>---------------------------------------------------------------------------</td>
<td>---------------</td>
<td>-----------</td>
</tr>
<tr>
<td>NOB DO half sat. [mgO₂/L]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>AAO DO half sat. [mgO₂/L]</td>
<td>0.0100</td>
<td>0.0100</td>
</tr>
<tr>
<td>Anoxic NO₃(→NO₂) half sat. [mgN/L]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Anoxic NO₃(→N₂) half sat. [mgN/L]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>Anoxic NO₂(→N₂) half sat. (mgN/L)</td>
<td>0.0100</td>
<td>0.0100</td>
</tr>
<tr>
<td>NH₃ nutrient half sat. [mgN/L]</td>
<td>0.0050</td>
<td>0.0050</td>
</tr>
<tr>
<td>PolyP half sat. [mgP/mgCOD]</td>
<td>0.0100</td>
<td>0.0100</td>
</tr>
<tr>
<td>VFA sequestration half sat. [mgCOD/L]</td>
<td>5.0000</td>
<td>0.0100</td>
</tr>
<tr>
<td>P uptake half sat. [mgP/L]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
<tr>
<td>P nutrient half sat. [mgP/L]</td>
<td>0.0010</td>
<td>0.0010</td>
</tr>
<tr>
<td>Autotroph CO₂ half sat. [mmol/L]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>H₂ low/high half sat. [mgCOD/L]</td>
<td>1.0000</td>
<td>1.0000</td>
</tr>
<tr>
<td>Propionic acetogens H₂ inhibition [mgCOD/L]</td>
<td>5.0000</td>
<td>5.0000</td>
</tr>
<tr>
<td>Synthesis anion/cation half sat. [meq/L]</td>
<td>0.0100</td>
<td>0.0100</td>
</tr>
</tbody>
</table>

**Common**

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biomass volatile fraction (VSS/TSS)</td>
<td>0.9200</td>
<td>0.9200</td>
</tr>
<tr>
<td>Endogenous residue volatile fraction (VSS/TSS)</td>
<td>0.9200</td>
<td>0.9200</td>
</tr>
<tr>
<td>N in endogenous residue [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in endogenous residue [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Endogenous residue COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
<tr>
<td>Particulate substrate COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.6000</td>
<td>1.6000</td>
</tr>
<tr>
<td>Particulate inert COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.6000</td>
<td>1.6000</td>
</tr>
<tr>
<td>Molecular weight of other anions [mg/mmol]</td>
<td>35.5000</td>
<td>35.5000</td>
</tr>
<tr>
<td>Molecular weight of other cations [mg/mmol]</td>
<td>39.1000</td>
<td>39.1000</td>
</tr>
</tbody>
</table>
### AOB

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield [mgCOD/mgN]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
<tr>
<td>AOB denite NO2 fraction as TEA [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Byproduct NH4 fraction to N2O [-]</td>
<td>0.0025</td>
<td>0.0025</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
</tbody>
</table>

### NOB

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield [mgCOD/mgN]</td>
<td>0.0900</td>
<td>0.0900</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
</tbody>
</table>

### AAO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield [mgCOD/mgN]</td>
<td>0.1140</td>
<td>0.1140</td>
</tr>
<tr>
<td>Nitrate production [mgN/mgBiomassCOD]</td>
<td>2.2800</td>
<td>2.2800</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
</tbody>
</table>
## OHO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield (aerobic) [-]</td>
<td>0.6660</td>
<td>0.3000</td>
</tr>
<tr>
<td>Yield (fermentation, low H2) [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Yield (fermentation, high H2) [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>H2 yield (fermentation low H2) [-]</td>
<td>0.3500</td>
<td>0.3500</td>
</tr>
<tr>
<td>H2 yield (fermentation high H2) [-]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Propionate yield (fermentation, low H2) [-]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Propionate yield (fermentation, high H2) [-]</td>
<td>0.7000</td>
<td>0.7000</td>
</tr>
<tr>
<td>CO2 yield (fermentation, low H2) [-]</td>
<td>0.7000</td>
<td>0.7000</td>
</tr>
<tr>
<td>CO2 yield (fermentation, high H2) [-]</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Endogenous fraction - aerobic [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>Endogenous fraction - anoxic [-]</td>
<td>0.1030</td>
<td>0.1030</td>
</tr>
<tr>
<td>Endogenous fraction - anaerobic [-]</td>
<td>0.1840</td>
<td>0.1840</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
<tr>
<td>Yield (anoxic) [-]</td>
<td>0.5400</td>
<td>0.5400</td>
</tr>
<tr>
<td>Yield propionic (aerobic) [-]</td>
<td>0.6400</td>
<td>0.6400</td>
</tr>
<tr>
<td>Yield propionic (anoxic) [-]</td>
<td>0.4600</td>
<td>0.4600</td>
</tr>
<tr>
<td>Yield acetic (aerobic) [-]</td>
<td>0.6000</td>
<td>0.6000</td>
</tr>
<tr>
<td>Yield acetic (anoxic) [-]</td>
<td>0.4300</td>
<td>0.4300</td>
</tr>
<tr>
<td>Yield methanol (aerobic) [-]</td>
<td>0.5000</td>
<td>0.5000</td>
</tr>
<tr>
<td>Adsorp. max. [-]</td>
<td>1.0000</td>
<td>1.0000</td>
</tr>
<tr>
<td>Max fraction to N2O at high FNA over nitrate [-]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>Max fraction to N2O at high FNA over nitrite [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
</tbody>
</table>
### Methylotrophs

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield (anoxic) [-]</td>
<td>0.4000</td>
<td>0.4000</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
<tr>
<td>Max fraction to N2O at high FNA over nitrate [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Max fraction to N2O at high FNA over nitrite [-]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
</tbody>
</table>

### PAO

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield (aerobic) [-]</td>
<td>0.6390</td>
<td>0.3000</td>
</tr>
<tr>
<td>Yield (anoxic) [-]</td>
<td>0.5200</td>
<td>0.5200</td>
</tr>
<tr>
<td>Aerobic P/PHA uptake [mgP/mgCOD]</td>
<td>0.9300</td>
<td>1.0500</td>
</tr>
<tr>
<td>Anoxic P/PHA uptake [mgP/mgCOD]</td>
<td>0.3500</td>
<td>0.3500</td>
</tr>
<tr>
<td>Yield of PHA on sequestration [-]</td>
<td>0.8890</td>
<td>0.9900</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>N in sol. inert [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous part. [-]</td>
<td>0.2500</td>
<td>0.2500</td>
</tr>
<tr>
<td>Inert fraction of endogenous sol. [-]</td>
<td>0.2000</td>
<td>0.2000</td>
</tr>
<tr>
<td>P/Ac release ratio [mgP/mgCOD]</td>
<td>0.5100</td>
<td>0.5100</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
<tr>
<td>Yield of low PP [-]</td>
<td>0.9400</td>
<td>0.9400</td>
</tr>
<tr>
<td>Mg to P mole ratio in polyphosphate [mmolMg/mmolP]</td>
<td>0.3000</td>
<td>0.3000</td>
</tr>
<tr>
<td>Cation to P mole ratio in polyphosphate [meq/mmolP]</td>
<td>0.1500</td>
<td>0.1500</td>
</tr>
<tr>
<td>Ca to P mole ratio in polyphosphate [mmolCa/mmolP]</td>
<td>0.0500</td>
<td>0.0500</td>
</tr>
<tr>
<td>Cation to P mole ratio in organic phosphate [meq/mmolP]</td>
<td>0.0100</td>
<td>0.0100</td>
</tr>
</tbody>
</table>
### Acetogens

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>H2 yield [-]</td>
<td>0.4000</td>
<td>0.4000</td>
</tr>
<tr>
<td>CO2 yield [-]</td>
<td>1.0000</td>
<td>1.0000</td>
</tr>
<tr>
<td>N in biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Fraction to endogenous residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
</tbody>
</table>

### Methanogens

<table>
<thead>
<tr>
<th>Name</th>
<th>Default</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetoclastic yield [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Methanol acetoclastic yield [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>H2-utilizing yield [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>Methanol H2-utilizing yield [-]</td>
<td>0.1000</td>
<td>0.1000</td>
</tr>
<tr>
<td>N in acetoclastic biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>N in H2-utilizing biomass [mgN/mgCOD]</td>
<td>0.0700</td>
<td>0.0700</td>
</tr>
<tr>
<td>P in acetoclastic biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>P in H2-utilizing biomass [mgP/mgCOD]</td>
<td>0.0220</td>
<td>0.0220</td>
</tr>
<tr>
<td>Acetoclastic fraction to endog. residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>H2-utilizing fraction to endog. residue [-]</td>
<td>0.0800</td>
<td>0.0800</td>
</tr>
<tr>
<td>Acetoclastic COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
<tr>
<td>H2-utilizing COD:VSS ratio [mgCOD/mgVSS]</td>
<td>1.4200</td>
<td>1.4200</td>
</tr>
</tbody>
</table>
Appendix E

Hydrodynamic shear force versus airflow

Airflow requirements were compared for a reactor with the following dimensions and dissolved oxygen:

<table>
<thead>
<tr>
<th>SWL</th>
<th>V</th>
<th>D</th>
<th>DO</th>
</tr>
</thead>
<tbody>
<tr>
<td>m</td>
<td>m³</td>
<td>m</td>
<td>mg/L</td>
</tr>
<tr>
<td>5</td>
<td>1571</td>
<td>20</td>
<td>6</td>
</tr>
</tbody>
</table>

The dimensions and dissolved oxygen were changed to see the sensitivity of airflow to design and operational parameters. Three airflows were calculated as a function of organic loading rate:

1) the airflow to achieve a minimum upflow superficial air velocity (i.e., hydrodynamic shear force); 2) the airflow for a heterotrophic reactor (i.e., no nitrification); and 3) the airflow for a nitrification reactor. The following equations were used to calculate the airflows.

For hydrodynamic shear force:

\[ USAV = \frac{OLR}{5} \]

\[ \text{Airflow} = USAV \cdot A_{reactor} \]

For the heterotrophic reactor (i.e., assuming 0.5 BOD/COD and 1.6 bCOD/BOD):

\[ P_{X,\text{bio}} = Y_H \cdot 1.6 \cdot 0.5 \cdot OLR \cdot V_{reactor} \cdot \frac{(1 + f_d \cdot b_H \cdot SRT)}{(1 + b_H \cdot SRT)} \]

\[ R_o = 1.6 \cdot 0.5 \cdot OLR \cdot V_{reactor} - 1.42 \cdot P_{X,\text{bio}} \]
For the nitrification reactor (i.e., assuming 8 COD/TN and 0.8 NOx-N/TN):

\[
P_{X,\text{bio}} = Y_H \cdot 1.6 \cdot 0.5 \cdot OLR \cdot V_{\text{reactor}} \cdot \frac{(1 + f_d \cdot b_H \cdot SRT)}{(1 + b_H \cdot SRT)} + Y_N \cdot 0.8 \cdot \frac{OLR}{8} \cdot V_{\text{reactor}} \cdot \frac{1}{(1 + b_N \cdot SRT)}
\]

\[
R_o = 1.6 \cdot 0.5 \cdot OLR \cdot V_{\text{reactor}} - 1.42 \cdot P_{X,\text{bio}} + 4.57 \cdot 0.8 \cdot \frac{OLR}{8} \cdot V_{\text{reactor}}
\]

For both reactors:

\[
SOTR = \frac{R_o}{\alpha \cdot F} \cdot \frac{C_{\infty,20}^*}{\left(\beta \cdot \frac{C_{st,20}^*}{C_{s,20}^*} \cdot \frac{P_b}{P_a} \cdot \frac{C_{\infty,20}^*}{C_{o,20}^*} - DO\right)} \cdot 1.024^{(20-T)}
\]

\[
\text{Air flow} = \frac{SOTR}{E/100 \cdot SWL \cdot 0.27}
\]

Other assumptions included:

<table>
<thead>
<tr>
<th>SRT</th>
<th>YH</th>
<th>bH</th>
<th>fd</th>
</tr>
</thead>
<tbody>
<tr>
<td>d</td>
<td>VSS/COD</td>
<td>1/d</td>
<td>-</td>
</tr>
<tr>
<td>30</td>
<td>0.45</td>
<td>0.12</td>
<td>0.15</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>COD/TN</th>
<th>YN</th>
<th>bN</th>
<th>NOx/TN</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSS/NOx-N</td>
<td>1/d</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>0.15</td>
<td>0.17</td>
<td>0.8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>α</th>
<th>F</th>
<th>β</th>
<th>T</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>C</td>
<td>%/m</td>
</tr>
<tr>
<td>0.65</td>
<td>0.9</td>
<td>0.95</td>
<td>20</td>
<td>4.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Df</th>
<th>de</th>
<th>Pa</th>
<th>C* s, 20</th>
<th>C* inf, 20</th>
<th>C* st</th>
<th>Pb</th>
</tr>
</thead>
<tbody>
<tr>
<td>m</td>
<td>m</td>
<td>mg/L</td>
<td>mg/L</td>
<td>mg/L</td>
<td>m</td>
<td></td>
</tr>
<tr>
<td>4.5</td>
<td>0.4</td>
<td>10.33</td>
<td>9.09</td>
<td>10.67</td>
<td>9.09</td>
<td>10.33</td>
</tr>
</tbody>
</table>
The following table depicts the results of airflow for all three cases (i.e., for the design and operational parameters presented at the beginning of the Appendix):

<table>
<thead>
<tr>
<th>Case OLR kg-COD/(m³∙d)</th>
<th>USAV Airflow m³/min</th>
<th>HET Airflow m³/min</th>
<th>AUT Airflow m³/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>0.5</td>
<td>19</td>
<td>25</td>
<td>43</td>
</tr>
<tr>
<td>1.0</td>
<td>38</td>
<td>50</td>
<td>86</td>
</tr>
<tr>
<td>1.5</td>
<td>57</td>
<td>75</td>
<td>128</td>
</tr>
<tr>
<td>2.0</td>
<td>75</td>
<td>100</td>
<td>171</td>
</tr>
<tr>
<td>2.5</td>
<td>94</td>
<td>124</td>
<td>214</td>
</tr>
<tr>
<td>3.0</td>
<td>113</td>
<td>149</td>
<td>257</td>
</tr>
<tr>
<td>3.5</td>
<td>132</td>
<td>174</td>
<td>300</td>
</tr>
<tr>
<td>4.0</td>
<td>151</td>
<td>199</td>
<td>343</td>
</tr>
<tr>
<td>4.5</td>
<td>170</td>
<td>224</td>
<td>385</td>
</tr>
<tr>
<td>5.0</td>
<td>188</td>
<td>249</td>
<td>428</td>
</tr>
<tr>
<td>5.5</td>
<td>207</td>
<td>274</td>
<td>471</td>
</tr>
<tr>
<td>6.0</td>
<td>226</td>
<td>299</td>
<td>514</td>
</tr>
<tr>
<td>6.5</td>
<td>245</td>
<td>323</td>
<td>557</td>
</tr>
<tr>
<td>7.0</td>
<td>264</td>
<td>348</td>
<td>600</td>
</tr>
<tr>
<td>7.5</td>
<td>283</td>
<td>373</td>
<td>642</td>
</tr>
<tr>
<td>8.0</td>
<td>302</td>
<td>398</td>
<td>685</td>
</tr>
<tr>
<td>8.5</td>
<td>320</td>
<td>423</td>
<td>728</td>
</tr>
<tr>
<td>9.0</td>
<td>339</td>
<td>448</td>
<td>771</td>
</tr>
<tr>
<td>9.5</td>
<td>358</td>
<td>473</td>
<td>814</td>
</tr>
<tr>
<td>10.0</td>
<td>377</td>
<td>498</td>
<td>857</td>
</tr>
<tr>
<td>10.5</td>
<td>396</td>
<td>523</td>
<td>899</td>
</tr>
<tr>
<td>11.0</td>
<td>415</td>
<td>547</td>
<td>942</td>
</tr>
<tr>
<td>11.5</td>
<td>434</td>
<td>572</td>
<td>985</td>
</tr>
<tr>
<td>12.0</td>
<td>452</td>
<td>597</td>
<td>1028</td>
</tr>
<tr>
<td>12.5</td>
<td>471</td>
<td>622</td>
<td>1071</td>
</tr>
<tr>
<td>13.0</td>
<td>490</td>
<td>647</td>
<td>1113</td>
</tr>
<tr>
<td>13.5</td>
<td>509</td>
<td>672</td>
<td>1156</td>
</tr>
<tr>
<td>14.0</td>
<td>528</td>
<td>697</td>
<td>1199</td>
</tr>
<tr>
<td>14.5</td>
<td>547</td>
<td>722</td>
<td>1242</td>
</tr>
<tr>
<td>15.0</td>
<td>565</td>
<td>746</td>
<td>1285</td>
</tr>
<tr>
<td>15.5</td>
<td>584</td>
<td>771</td>
<td>1328</td>
</tr>
<tr>
<td>16.0</td>
<td>603</td>
<td>796</td>
<td>1370</td>
</tr>
<tr>
<td>16.5</td>
<td>622</td>
<td>821</td>
<td>1413</td>
</tr>
<tr>
<td>17.0</td>
<td>641</td>
<td>846</td>
<td>1456</td>
</tr>
<tr>
<td>17.5</td>
<td>660</td>
<td>871</td>
<td>1499</td>
</tr>
<tr>
<td>18.0</td>
<td>679</td>
<td>896</td>
<td>1542</td>
</tr>
<tr>
<td>18.5</td>
<td>697</td>
<td>921</td>
<td>1585</td>
</tr>
<tr>
<td>19.0</td>
<td>716</td>
<td>946</td>
<td>1627</td>
</tr>
<tr>
<td>19.5</td>
<td>735</td>
<td>970</td>
<td>1670</td>
</tr>
<tr>
<td>20.0</td>
<td>754</td>
<td>995</td>
<td>1713</td>
</tr>
</tbody>
</table>