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Microwaving human faecal sludge as a viable sanitation technology option for treatment and value recovery – a critical review

Authors

• Oluwasola O.D. Afolabi
• M. Sohail

Affiliations

School of Civil and Building Engineering, Loughborough University
Loughborough, LE11 3TU, UK

*Corresponding author

E-mail address: o.o.d.afolabi2@lboro.ac.uk

Highlights

• Human faecal sludge is not a waste but an organic-rich resourceful material
• Moisture content of human faecal sludge makes them suitable for dielectric heating
• Recovery of pathogen-free value-added products such as chars, fertilizers is feasible
• Value-added products recovered can provide financial benefit
• Parameters for scaling the microwave technology for sanitation purposes are proposed
Abstract

The prolonged challenges and terrible consequences of poor sanitation, especially in developing economies, call for the exploration of new sustainable sanitation technologies. Such technologies must be: capable of effectively treating human faecal wastes without any health or environmental impacts; scalable to address rapid increases in population and urbanization; capable of meeting environmental regulations and standards for faecal management; and competitive with existing strategies. Further and importantly, despite its noxiousness and pathogenic load, the chemical composition of human faecal sludge indicates that it could be considered a potentially valuable, nutrient-rich renewable resource, rather than a problematic waste product. New approaches to faecal sludge management must consequently seek to incorporate a ‘valuable resource recovery’ approach, compatible with stringent treatment requirements.

This review intends to advance the understanding of human faecal sludge as a sustainable organic-rich resource that is typically high in moisture (up to 97 per cent), making it a suitable candidate for dielectric heating, i.e. microwave irradiation, to promote faecal treatment, while also recovering value-added products such as ammonia liquor concentrate (suitable for fertilizers) and chars (suitable for fuel) – which can provide an economic base to sustain the technology. Additionally, microwaving human faecal sludge represents a thermally effective approach that can destroy pathogens, eradicate the foul odour associated human faecal sludge, while also preventing hazardous product formations and/or emissions, aside from other benefits such as improved dewaterability and heavy metals recovery. Key technological parameters crucial for scaling the technology as a complementary solution to the challenges of onsite sanitation are also discussed.

Keywords

Human faecal sludge, microwave, sanitation, waste management, resource recovery, low- and middle-income countries
1. Introduction

Despite numerous longstanding international programmes and complementary strategies aimed at addressing poor sanitation in at-risk countries, the statistics for people without improved sanitation facilities and the associated health risks and effects are still alarming: 2.4 billion people currently lack access to improved sanitation facilities, while 1.1 billion still practice open defecation (WHO/UNICEF, 2015) (see Fig. 1).

**Fig. 1. Global distribution of people (numbers in millions by region) without access to improved sanitation facilities in 2015 [Source: WHO/UNICEF, 2015]**

The consequences are devastating: poor sanitation and contaminated drinking water resources cause more than 80 per cent of all disease in the developing world (WHO, 2003; CDC, 2011). An estimated 1.8 billion people consume faecal-contaminated water, and this increases the health risk of water-borne diseases – such as diarrhoea – which kill 700,000 infants per year (UNICEF-IGME, 2012). Apart from the health impacts, poor sanitation also has pervasive societal effects, such as increasing the rate of school drop-outs for young women and girls who cannot attend school without the adequate sanitation facilities they need to manage their personal hygiene during menstruation (WHO/UNICEF, 2004). Furthermore, nursing the sick that contract diseases due to poor sanitation adds an additional financial strain on many already living below the US$1.25 per day poverty line, while also reducing their ability to earn incomes, thereby encouraging a cycle of continuous poverty.
Apart from the non-technical issues – such as lack of a regulatory framework, sanitation-related institutions, capacities and financial constraints – most approaches to onsite sanitation in economically disadvantaged regions may be viewed as being labour intensive, crude and ineffective. Such approaches fail to address basic structural, environmental and ecological needs (Eckhoff and Wood, 2011). Currently, open defecation and manually emptying of human faecal sludge (HFS) slurry-pits, with the accompanying pumping, draining, storage, transportation and disposal, is still a widespread practice. Such activities create hazardous exposure to pathogens for all concerned, and often lead to uncontrolled and indiscriminate disposal into drains, canals and open places, especially in developing countries. Indiscriminate disposal sometimes take place near points of (water) abstraction or near slums and shantytowns of economically deprived communities due to poor road networks, traffic congestion and a lack of properly managed dumps/landfill sites (Strauss and Montangero, 2002). Such practices further aggravate the health impacts of exposure to untreated HFS and pollution of the environment. For millions of people across the developing world, restricted access to adequate sanitation, hygiene and potable water affects their nutritional status, health and quality of life, and is ultimately reflected in the reduced life expectancies characteristic of communities that are impacted in this way.

The effects of poor sanitation demand the introduction of complementary approaches, which should be strategically positioned to resolve current impacts and mitigate those induced by increasing urbanization and populations. Such approaches need to embody effective, affordable and sustainable technologies, which can substitute/complement current approaches to onsite sanitation. A review of existing technologies, such as the widely used biological processes and incineration, finds that they are difficult to implement in dispersed rural settlements and poorly developed urban settings with high occupation rates (slums). Large construction footprints, long conversion times, greenhouse gas emissions, along with air pollutants, all raise further barriers to their transfer/adoption in economically deprived areas. Consequently, there is an enduring need to develop better technologies, or remodel and optimize existing ones that are sustainable and effective in managing human faecal sludge. Most importantly, such developments must be viable and able to be scaled to meet the needs of an increasing population that is undergoing rapid urbanization.

Furthermore and also importantly, a large proportion of the nutrients in food consumed are present in urine and faeces, and hence HFS contains valuable resources that include, but are not limited to: water, nutrients, urea, minerals, salts and energy. Two primary components that can be viably recovered from HFS are plant nutrients (primarily nitrogen [N], phosphorus [P], potassium [K]) and energy [carbon, C]). Nevertheless, the pathogenic content of HFS presents a
serious challenge, both for treatment and resource-recovery purposes. Additionally, HFS may also contain residues of many complex engineered chemicals/synthetic products – hormones, antibiotics and food additives, for example. Some of these compounds pose further environmental risks due to their poor biodegradability. The challenge, therefore, is to develop novel technological ways and means necessary to provide safe and affordable sanitation to 2.4 billion people in a manner that is pleasing to use, publically acceptable and that effectively removes human waste from the environment, while also recovering recyclable valuables from it. Ideally, this is to be achieved without additional financial burden or the need for piped water or sewerage systems, which are not present/readily available in regions affected by poor sanitation. This need forms the basis for this review, which seeks to:

- illustrate HFS as a ‘resourceful’ material that can be beneficially exploited for the realization of its intrinsic value (to provide energy and plant fertilizer, for instance);
- advance the knowledge of microwave suitability as an alternative sanitation technology for processing HFS – to convert it into a safe, pathogen-free material, while also recovering inherent value; and
- enumerate parameters and other related factors crucial for the scaling of the technology for onsite sanitation/ faecal management purposes.

2. Human faecal sludge characterization

HFS, in the context of onsite sanitation, generally encompasses the slurry/semi-solids mixture of fresh, unprocessed or partially digested excreta (a mixture of urine and faeces only), along with associated sanitary cleansing products typically found at and in sanitation facilities such as pit latrines, wash blocks, mobile toilets, septic tanks, aqua privies, dry toilets and open-defecation sites. It may also contain blackwater, with or without greywater, and makes up the largest amount of bodily waste generated daily (Franceys et al, 1992). HFS is highly heterogeneous and variable in content, consistency, quantity, composition and concentration (Hemkend-reis et al, 2008). Fig. 2 shows the characteristic composition of HFS. It is characterized as a complex mixture of water; pathogenic microorganisms (viruses, bacteria, helminths and protozoa); nutrients in the form of undigested macromolecular organics (i.e. protein, lipids and polysaccharides); and inorganics, ash and minerals (Franceys et al, 1992; Putman, 1971; Wignarajah et al, 2006). The presence of exogenous compounds, including radioisotopes, antibiotics, food additives and heavy metals, has also been reported in HFS (Schouwa et al, 2002; Titley et al, 2000; Vinnerås et al, 2006).
Fresh HFS is typically brown in colour and tends towards dark brown, depending on length of storage. The Bristol Stool Chart described by Heaton and Lewis (1997), which ranges from Type 1 through to Type 7, is widely used as a linear scale to characterize the texture, forms and variation of HFS based on visual appearance. Foul odour characterizes HFS due to the presence of nitrogenous benzopyrrole compounds, notably indole and skatole, hydrogen sulphides, methyl sulphides and other sulphur-containing compounds (Moore et al, 1987; Sato et al, 2002). This has an instantaneous, unpleasant impact, causing considerable distress to the public and workers, especially during handling, storage and/or transportation. While the toxicological effects of foul odours associated with HFS are yet to be proved, there have been reported cases of illness/symptoms associated with exposure to malodorous sludges (Chrostowki, and Foster, 2003; Cain, and Cometto-Muniz, 2004).

The moisture content in human faeces is between 63 and 86 per cent w/w (Rose et al., 2015), while urine is mainly composed of water: between 93 and 96 per cent w/w (Polprasert, 1995). The addition of flush water, anal cleansing water and/or other sanitary products affects the moisture-to-total solids loading in resultant HFS. Up to 97 per cent moisture content – similar to those obtained in primary sewage sludge (commonly used as surrogate for faecal sludge for treatment purposes) – has been reported in HFS (Heinss et al, 1998; Tchobanoglous et al, 2003; Niwagaba et al, 2014). The pH of faecal sludge water generally ranges between 5.0 and 9.3 (Mai et al, 2009; Kengne et al 2011), influenced by dietary fibre in faecal matter (Newark and Lupton, 1990) and potentially by composition of sanitary products and the buffering effects of urine and flush water.

Untreated HFS is hazardous due to the potential presence of pathogenic bacteria, viruses, parasitic protozoa and helminthic nematodes. It is the faecal matter in HFS that contains most of these pathogens, as urine from healthy humans is generally sterile (WHO, 2006). Human faeces may contain a pathogenic load in the range 10^5–10^10 faecal coliforms per 100 ml (Von Spelling, 2007) and up to 60,000/L of helminth eggs (Heinss et al, 1998). To enable safe handling and reuse of treated HFS value-added products, the levels of *E.coli* and/or helminth eggs in treated HFS are used as health/safety pathogenic indicators. These must be: for *E.coli* (number per 100ml of treated waste) <1000/g total solids, while for helminth eggs (number per gram total solids or per litre) <1/g total solids (WHO, 2006).

Analyses of faecal solids reveal they are composed of 84–93 per cent organics, with the remainder as inorganics fractions (Feachem et al, 1978; Nwaneri et al, 2008; Bai and Wang, 2011). The inorganics are composed mainly of phosphates of calcium, iron and internal organs’
secretions (Guyton and Hall, 2000), while the organic fraction is composed of bacteria debris, typically 10–30 per cent (Ganong, 1983; Wignarajah et al, 2006) and up to 54 per cent (Stephen, and Cummings, 1980); and undigested protein – between 2 and 25 per cent, with up 50 per cent contribution from dead bacteria debris (Canfield et al, 1963; Volk and Rummel, 1987). Undigested fats are in the range 2–15 per cent (Kien et al, 1981; Chen et al, 1998; Wierdsma et al, 2011) and up to 25 per cent (Wignarajah et al, 2006); carbohydrates 10–30 per cent (Ganong, 1983); and minerals – mainly calcium (Ca), potassium, (K) and phosphorus (P) – in the range 5–8 per cent (Wignarajah et al, 2006).

These percentage weight distributions are largely affected by diet. Undigested proteins in faecal matter are mainly nitrogenous compounds from bacteria biota, nucleic acids and mucus secreted from intestinal walls (Bender and Bender, 1997). Fatty acids, waxes and phosphoglycerides make up the fat content in faecal matter, and are derived from undigested dietary fat intake, bacteria and fat in shredded epithelial cells (Guyton and Hall, 2000; Rose et al, 2015), while faecal carbohydrates are mainly composed of undigested cellulose, fibres and pentosan (Canfield et al, 1963). Undigested fibres make up to 25 per cent of the carbohydrates fraction (Volk and Rummel, 1987). Organics in urine solids range from 65 to 85 per cent (Strauss, 1985) and are predominantly composed of urea (Putman, 1971). The urine liquid fraction is composed mainly of water, organic acids (uric acid, citric acids), electrolytes, solubilized metabolites, hormones, vitamins, ammonium salts, nitrogenous compounds and inorganic salts (Putman, 1971; Polprasert, 1995).

Elemental characterization of dried faecal matter fraction in HFS reveals it is composed of 4.5 per cent Ca; 3–5.4 per cent P and 1–2.5 per cent K (Feachem et al, 1978), while dried urine is composed of 14–18 per cent N; 13 per cent C; 3.1 per cent P and 3.7 per cent K (Strauss, 1985). Faecal fractions generally contain more heavy metals, including zinc (Zn), copper (Cu), nickel (Ni), cadmium (Cd), lead (Pb) and mercury (Hg) (Schouwa et al, 2002). Generally, the concentrations of heavy metals reportedly found in faecal sludge from different studies are relatively low and in most cases less than 2mg/cap/day – with the exception of Zn, which can be up to 16mg/cap/day (Hansen and Tjell, 1979; Eastwood et al, 1984; Schouwa et al, 2002; Vinnerås et al, 2006).

Generation rates of HFS vary largely and depend on many factors such as gender, age, occupation, geographical location and climate, body weight, dietary intake and water consumption (Franceys et al, 1992; Rose et al, 2015). Direct measurement is primarily employed in studies to estimate generation rates. These find a range of 130–520g/cap/day of faeces
generated in developing countries and 100–200 g/cap/day for European and North American countries (Feachem et al, 1983). Differences in generation rates across the regions concerned are largely due to the nature of diet, as developing countries consume a more fibre-based diet, while and European and North American countries consume a more protein-based diet (Franceys et al, 1992; Wignarajah et al, 2006; Eastwood et al, 1984). The amount of urine generated is also largely dependent on the water consumption rate, temperature and humidity defining the geographical location considered. Generally, urine-generation rates range between 0.6 and 2.1 L/cap/day (Wydeven and Golub 1991; Del Porto and Steinfeld, 1999; Schouwa et al, 2002). Additionally, stooling rates/frequency of urination, waste streams due to sanitary products, and water inputs due to flushing and/or anal cleansing (which are largely dependent on occupation and behavioural practices) affect daily generation rates for HFS. Generation rates of these additional waste streams are also summarized in Fig. 2. For faecal treatment design purposes, one review recommended an average of 128 g/cap/day and 1.42 L/cap/day of faecal and urine generation rates (Rose et al, 2015), as use of flush water volume and other sanitary-related products could be managed or controlled.
Fig. 2. Characteristic composition of HFS (extracted from Heinss et al, 1998; Schouwa et al, 2002; Wignarajah et al, 2006; Niwagaba et al, 2014; Rose et al, 2015)

**Elemental composition**
- Category 1: Present in significant amounts: carbon (C), oxygen (O), hydrogen (H), nitrogen (N), phosphorus (P), potassium (K) and sulphur (S)
- Category 2: Present in smaller but significant amounts: calcium (Ca), magnesium (Mg) and zinc (Zn)
- Category 3: Trace elements such as copper (Cu), nickel (Ni), cadmium (Cd), lead (Pb), mercury (Hg)

**Pathogenic composition**
- Contains bacteria, viruses, helminths, protozoa
- Pathogenic loads range
  - 2,500–60,000 helminth eggs/L
  - $10^5$–$10^{10}$ faecal coliforms/100ml

**Chemical composition (dried basis)**
- Fat content/lipids: 2–15%
- Protein (and other nitrogenous materials): 2–25%
- Carbohydrates: 10–30%
- Minerals (K, Ca & P): 5–8%
- Bacterial debris: 10–54%
- Urea fraction in urine: over 50%
- Calorific value
  - Faeces: 17–21MJ/kg
  - Urine: 7.1 MJ/kg

**Physical characteristics**
- Color: Brown
- Sensory assessment: Foul
- Moisture content (MC): 95–97%
- Total solids (TS): 3–5%
- Volatile solids of TS: 45–73%
- Fixed solids of TS: 27–55%
- pH: 5–9
- Electricity conductivity
  - Urine: 160ms/cm

**Organic composition**
- COD: 20–50 g/L
- BOD: 6–7.6 g/L
- TP: 150–450 mg/L
- TN: 190–300 mg/L
- TKN: 1–3.4 g/L
- Nitrates: 0.2–21 mg/L
- TOC: 8–10g/l

**Generation rates**
- Faeces: 130g–530 g/cap/day
- Urine: 0.6–2.1 L/cap/day
- Anal cleansing water: 0.35–3L/wash
- Flush water: 1–12L/flush
- Toilet paper: 11.7–36g/cap/day
- Menstrual pads and flow: 34g/cap/day
- Other sanitary items: 0.2g/cap/day
- Stooling rate: 0.2–2.5/cap/day
- Urine frequency: 5.4–8 /cap/day
2.1 Assessment of intrinsic value recoverable from HFS

HFS has been explored extensively for agricultural use (Cofie et al, 2005; Cofie et al, 2006; Yadav et al, 2010). Consider, for example, Terra preta do Índio, also called Amazonian Dark Earth (ADE) – highly fertile and productive black soils found in the Amazon Basin region, which are reported to result from the prolonged use of soil ameliorants derived from human waste (Lehmann et al, 2003; Yoshida and Antal, 2009). Direct excretion of urine and faeces onto agricultural land in Thailand to supplement fertility has also been reported (Schouwa et al, 2002). At the same time, the direct application of urine for horticulture is a common practise that, in Nepal, has been scaled up so that diverted urine stored in urine banks may be either applied directly to soil, used in aquaculture or used to produce struvite (a phosphorous-rich, powdery compound derived from the addition of magnesium to urine banks), which is subsequently applied as fertilizer (Etter et al, 2011).

Essentially the key plant nutrients in HFS that are vital from the resource-recovery perspective are nitrogen (N), phosphorus (P) and potassium (K). Table 1 provides ranges of daily generation rates of nutrients recoverable from the urine and faecal components of HFS. Of the faecal and urine fractions of HFS, urine contains the largest amount of N (90 per cent); P (50–65 per cent) and K (50–80 per cent) (Heinonen-Tanski and van Wijk-Sijbesma, 2005). P and K in urine are excreted in ionic forms and in plant-available forms that are the same as those found in chemical fertilizers (Kirchmann and Pettersson 1995). Studies conducted in Sweden indicate that on an yearly basis nutrients recoverable per capita in urine are: 2.5–4.3kg N, 0.4–1.0kg P and 0.9–1.0kg K, while those in the faecal component are: 550g N, 183g P and 365g K (Lentner and Wink, 1981; Jonsson et al, 2005; Vinnerås et al, 2006). This means the plant nutrients in the urine and faecal components add up to 4.8kg N, 1.1kg P and 1.4kg K per capita per year (Jonsson et al, 2005; Vinnerås et al, 2006); this is enough to meet the fertilizer requirements for about a quarter ton of cereal, for example (Drangert, 1998).
Table 1 Daily generation loading of key plant nutrients in HFS

<table>
<thead>
<tr>
<th>Plants nutrients</th>
<th>Faeces Ranges of generation rates (g/cap/day) (wet basis)</th>
<th>Urine Ranges of generation rates (g/cap/day) (wet basis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total N</td>
<td>0.3–1.51</td>
<td>2–351</td>
</tr>
<tr>
<td>Total P</td>
<td>0.352–2.73</td>
<td>0.45–1.38</td>
</tr>
<tr>
<td>Total K</td>
<td>0.24–2.52</td>
<td>0.783–2.874</td>
</tr>
<tr>
<td>Na</td>
<td>0.3–4.15</td>
<td>0.088–4.531</td>
</tr>
<tr>
<td>Ca</td>
<td>0.13–3.66</td>
<td>0.129–0.531</td>
</tr>
<tr>
<td>Mg</td>
<td>0.154–0.354</td>
<td>0.194–0.214</td>
</tr>
</tbody>
</table>

1Extracted from Jonsson et al, 2005  2Extracted from Vinnerås et al, 2006
3Extracted from Wignarajah et al, 2006  4Extracted from Calloway and Margen, 1971
5Extracted from Eastwood et al, 1984  6Extracted from Chaggu, 2004
7Extracted from Rose et al, 2015  8Extracted from Borawski et al, 2008
9Extracted from Snyder et al, 1975

From an energy-recovery perspective, recovery of constituent carbon (C) in HFS is essential. Faecal and urine components of HFS contain 44–55 per cent and 11–17 per cent C respectively (Feachem et al, 1978; Strauss, 1985). More than 7g/cap can be recovered based on daily loadings estimation (Snyder et al, 1975). Similarly, the volatile solids, which provide an indication of combustibility, range between 75 and 85 per cent in urine and up to 92 per cent in faeces (Fry and Merrill, 1973). When combined in HFS, however, this figure ranges between 45 and 73 per cent (Niwagaba et al, 2014).

Carbon content in any material correlates with its calorific value. The use of bomb calorimeters, proximate analysis (volatile matter, fixed solids and moisture content determination by standard procedure) or elemental analyses (determination of carbon, hydrogen and nitrogen contents) of materials are sometimes used to predict the amount of energy that can be recovered from such material (Cordero et al, 2001; Friedl et al, 2005). Based on these, one study predicted the average calorific value of dried urine and dried faeces to be 7.1MJ/kg and 17MJ/kg respectively (Rose et al, 2015). Another reported the calorific values of faecal sludge in three African countries to range between 16 and 19MJ/kg (Muspratt et al, 2014). For biogas recovery through anaerobic digestion, for example, or for other related biofuels recovery, chemical oxygen demand (COD) values of HFS, which range between 20 and 50g/L (Niwagaba et al, 2014) at 1.6kg/cap/day (Vinnerås et al, 2006), suggest they can be suitable substrates. However, direct use of HFS for anaerobic digestion is not recommended and requires careful consideration, as the levels of ammonia, organics (urea for example), heavy metals and sulphides present in HFS are key
inhibitors of principal microorganisms (methanogens in particular) of the anaerobic digestion process (Chen et al, 2008).

In essence, a good percentage of diet consumed is locked as an energy resource and plant nutrients in HFS. Using an appropriate technological technique – like the microwave processing reviewed in this paper – these resources/nutrients can be beneficially recovered for other applications, without any of the health risks or environmental impacts that hamper their direct use.

2.2 Case for novel technological integration with onsite sanitation systems

From a health perspective, direct use or exposure to untreated HFS is always hazardous (due to its pathogenic load) and represents a risk to public health. Any treatment method must address such risks and conform to World Health Organization (WHO) guidelines on E.coli and helminth eggs, due to their prevalence in both onsite sanitation and conventional wastewater treatment facilities and their propensity to end up in the biosolids generated by wastewater treatment (WHO, 2006). Over the years, few onsite sanitation solutions used in developing countries have comprehensively addressed pathogen kill and odour eradication while treating HFS. Composting, for example, is characterized by significant pathogenic residues due to the low temperatures (<50°C) involved (Schoenning and Stenstroem, 2004; Spuhler, 2011). Direct exposure to faecal matter during the composting process also represents a health risk. Other biological processes – such as settling tank/sedimentation beds, unplanted natural air-drying beds and wetlands – are ineffective, as they also cannot generate pathogen-free stabilized biosolids. Odour nuisance, parasitic helminths (which tend to concentrate in settled or floating solids (Jimenez, 2007)) and E. coli, for example, characterize the end products of these processes (WHO, 2006). Anaerobic digestion technologies require significant construction investments, can take up to three weeks to process human waste and still do not ensure complete pathogen destruction. The highest temperature range of anaerobic digestion, i.e. the thermophilic range, is 45–60°C (Demirbas et al, 2011); this is too low for pathogenic inactivation. Freeze/thawing technology, which damages pathogenic cells via intracellular freezing and osmotic effects (Mazur, 1986), has recorded little success in terms of pathogen deactivation, as thawed sludge falls short of WHO treatment standards (Sanin et al, 1994). Also, the process is highly selective and while economic for poor cold regions, is impractical for regions characterized by relatively high ambient temperatures. The process additionally requires high-energy inputs due to the inevitable low temperature
requirement for freezing. Thermal treatment, such as incineration, is capital intensive and generates dioxins and other toxic air pollutants, which pose risks to public health (US EPA, 2003; Rulkens, 2008). Furthermore, such processes are limited to dry substrate as the starting material, requiring high-energy inputs with significant greenhouse gas emissions (Demirbas et al, 2011). Other technologies such as ultrasonication (Show et al, 2007), hydrodynamic cavitation (Lee and Han, 2013); mechanical treatment (Nah et al, 2000); oxidation by ozone (Yeom et al, 2002); alkali pretreatment (Mouneimne et al, 2003); or a combination of technologies (Barienbruch and Kopplow, 2003; Eskicioglu et al, 2008b; Zupanc et al, 2013; Yeneneh et al, 2015; Doğruel, and & Özgen, 2016) have also been reported. While most of these technologies have been extensively explored as sludge pre-treatment processes, their potential use and/or integration into onsite sanitation systems is still evolving.

In summary, most of these current approaches are difficult to implement in dispersed rural settlements and poorly developed urban settings with high-occupation rates (slums). The large construction footprints, long conversion times, greenhouse gas emissions, along with air pollutants, all raise further barriers to their transfer/adoption in economically deprived areas. Consequently, there is an enduring need to develop better technologies, or to remodel and optimize existing ones that are sustainable and effective at managing HFS. Such developments must be viable for adoption and transfer in low- and middle-income countries.

3. The microwave technology

Microwave technology is an acknowledged and widely accepted heating source, and has been used for diverse applications. Developed primarily for communications and some areas of food processing, microwaves are now extensively explored for digestion of environmental samples into ionic forms for quantitative analyses (Islam, and Weil, 1998); and accelerated synthesis of materials to generate and produce new and valuable intermediates/compounds, and nanostructure materials with special functionalities (Ifrah et al, 2007; Zhenyu et al, 2007; Hu et al, 2009; Wang et al, 2011). This technology has also been used to recover valuable gaseous hydrocarbons from dried sewage sludge (Menendez et al, 2002; Sobhy and Chaouki, 2010). The feasibility of microwave technology to achieve these results has been attributed to its characteristic potential to selectively heat materials and alter (suppress and/or activate novel) reaction pathways, and accelerate overall reaction rates through the induction of rapid volumetric heating at the molecular level (Clark et al, 2000; Sobhy and Chaouki, 2010). The net effects of
this include reduction on processing times, energy and associated costs. Microwave irradiation also has the capacity to diminish hazardous product formation and prevent emission of fugitive or greenhouse gases, hence making it environmentally friendly.

In essence, microwave technology has significant potential as a viable and efficient heating source for the treatment and valorization of bio-based wastes/materials such as HFS. Furthermore and most importantly, technological familiarization and established public acceptance of microwave technology as a domestic (homes and offices) and industrial appliance, both in developed and developing regions, may promote its adoption and transfer into existing onsite sanitation systems — either as a standalone unit or integrated with a network of decentralized systems. Despite its potential, however, knowledge or application of this technology for treating and processing human excreta or HFS\(^1\) for onsite sanitation is largely non-existent. This review partly intends to drive interest in the potential of microwave technology for processing HFS at household, small networked decentralized onsite sanitation systems and other related facilities, and as a technological option that can be used for mobile processing of HFS.

3.1 Principles of the microwave technology

Within the electromagnetic spectrum, microwaves are non-ionizing radiation that occupy the wavelengths from 1mm to 1m, with a corresponding frequency range of 0.3GHz–300GHz (Shuntaro and Jun-Ichi, 2011). Microwave heating applied for industrial, medical and scientific applications use mainly two microwave frequencies, 0.915GHz and 2.45GHz (Bradshaw et al, 1998). Domestic ovens, for example, use the 2.45GHz frequency to heat. At this frequency, the energy of the microwave photon is 0.0016eV; this is too low to break chemical bonds such as H-OH, which is 5.2eV, and hence cannot ionize human living tissue, as compared with the ~33eV energy released per ionization radiation of x-rays/gamma rays (Kappe, 2004; Yin, 2012). This means microwaves are not ionizing radiation, cannot induce chemical reactions and nor can they affect users of the technology within necessary safety limits.

\(^1\) This does not apply to primary sewage sludge obtained in wastewater treatment plants, as this is different in form and physicochemical compositions to fresh human excreta or HFS in that it has undergone significant decomposition and degradation during transit in sewers, and may contain non-domestic waste streams. However, this doesn't preclude microwave suitability for primary sewage sludge treatment or valorization.
Microwave systems have three main integrated features: microwave generation, transmission and application (See Fig. 3). Microwaves are usually generated in a vacuum tube, commonly referred to as a magnetron or a magnetron tube, and coupled for transmission through a waveguide (hollow tubes) to the point of application via microwave (single/multi-mode cavities) applicators (Thostenson and Chou, 1999). For heat to be generated within a material by microwaves, it must be partially transparent to microwave radiation while absorbing the microwave energy. The ability of a material to absorb microwave energy is governed by its dielectric properties: dielectric constant ($\varepsilon'$) and dielectric loss ($\varepsilon''$). The dielectric constant quantifies its capacitive property (i.e. the ability of the dipolar molecule of the material to be polarized by an electric field), while dielectric loss reflects its conductive ability (i.e. the efficiency of converting incident microwave energy to heat within the material) (Bradshaw et al, 1998; Thostenson and Chou, 1999). Both properties are related by the complex dielectric constant ($\varepsilon^*$); this is used to describe the total dielectric properties of any material and is expressed as:

$$\varepsilon^* = \varepsilon' \pm j\varepsilon''$$  \hspace{1cm} \text{Equation 1}$$

Where the real part, $\varepsilon'$, represents the ability of the material to be polarized by an external electric field; and the imaginary part, $j\varepsilon''$, is the effective loss which quantifies the efficiency with which the microwave energy is converted to heat (Venkatesh and Raghavan, 2004).

Fig. 3. Schematic features of a microwave system
Tangent loss, also called the dissipation factor of materials, tan δ, is the ratio of the dielectric loss, $\varepsilon''$, of heated material to its dielectric constant, $\varepsilon'$. Tan δ reflects the dielectric response of any materials and is used to describe the overall efficiency of any material to absorb microwave energy and convert it to heat at a specific frequency (Clark et al, 2000). It is used to characterize the microwave-absorbing potentials of materials such as: transparent/insulators (i.e. materials not sensitive to microwave energy, as it can pass through with little or no attenuation, e.g. glass, Teflon, quartz, ceramic); reflecting/opaque materials (i.e. materials that do not allow microwaves to penetrate through them and react by reflecting microwaves (e.g. metals and other conductors); and receptors/dielectrics (i.e. good absorbers of microwaves, depending on their dielectric properties (e.g. activated carbon, water, electrolytes) (Larhed et al, 2002; Kappe, 2004). Generally, materials with tan δ > 0.5 are referred to as good microwave absorbers, while those with tan δ between 0.1 and 0.5 and those < 0.1 are considered to be medium and low-microwave absorbers, respectively (Yin, 2012). However, a low tan δ material can be blended with materials with a high tan δ to improve its overall microwave absorptive capacity, as suggested in previous studies (Menendez et al, 2002; Sobhy and Chaouki, 2010).

4. **Technological feasibility of microwave heating for HFS processing**

4.1 **Suitability of microwave heating for processing HFS**

The theory behind the adoption of microwave heating for processing HFS is based on the concept that water, which constitutes up to 97 per cent w/w in HFS, efficiently couples with microwaves and this electromagnetic interaction causes dielectric heating. In other words, the electromagnetic copulation of water and other molecular dipolar organics in HFS with incident microwaves triggers dielectric heating that occurs at the molecular level. This can achieve both pathogenic deactivation and value recovery from HFS at optimal process conditions – such as microwave power, temperature and residence time of processing. Two main dielectric heating mechanisms explain this feasibility: thermal effects due to dipolar polarization/rotation and athermal effects triggered by ionic conduction/migration as shown in Fig 4 (Kappe, 2004; Yin, 2012).
Dipolar polarization is the primary heat-induction mechanism of microwave dielectric heating, which results from the direct coupling of molecular dipoles with electromagnetic microwaves. Dipolar materials such as water are highly sensitive to electric fields, as their local charges tend to move in response to an applied electric field. Dipolar polarization results from the interaction of incident microwave irradiation with molecular dipoles, causing the dipoles to respond by rotating in order to realign in the direction of the applied electromagnetic field. As the dipole re-orientates to align itself with the field, the field is already changing, creating a phase difference between the orientation of the field and the dipole. This phase difference causes energy to be lost from the dipoles through random collisions/frictions at a frequency of 2.45GHz (i.e. occurring at 2.45 billion times per second), causing dielectric heating. In other words, dipole polarization lags behind an applied electric field due to the internal forces and/or bond resistance of realigning dipolar molecules, and this lag results in molecular friction or dielectric loss – characterized by energy loss dissipated in the form of heat (Bradshaw et al, 1998; Clark et al, 2000). Athermal effects, i.e. those not associated with temperature increase, have been attributed to the net effect of the continuous realignment of polarized dipoles to the electric field, and ionic migration – which occurs when an applied electrical field causes dissolved ions of positive and negative charges of heated dipolar materials to move towards oppositely charged regions of the field. This results in net collisions of dipoles and possible weakening of the hydrogen bonds within water molecules of heated material (Kappe, 2004; Venkatesh and Raghavan, 2004).

Fig. 4. Microwave dielectric heating mechanisms (Source: Yin, 2012)
4.2 Potential effects of microwave heating interaction with HFS

4.2.1 Odour eradication through thermal solubilization

Microwave heating is capable of eradicating foul odour associated with HFS through thermolytic solubilization. Thermal hydrolysis during microwave heating can solubilize organic components of HFS, including those associated with foul odour, trapping them in the aqueous phase. As described in Section 2.0, the primary organics of HFS are carbohydrates, proteins and lipids. During microwave heating, these macromolecular compounds are hydrolyzed into shorter-chain intermediate monomers: carbohydrates into reducing sugars; proteins into saturated and unsaturated acids such as amino-acids, ammonia and carbon dioxide; while lipids are broken down into short-chain fatty acids such as palmitic acids and oleic acids (Lian-hai et al, 2006; Wilson and Novak, 2009). Studies have demonstrated this thermolytic solubilization capacity of microwave heating. Ahn et al, (2009) applied microwave heating at 700W for 15 minutes on sewage sludge and observed increases in concentrations of soluble carbohydrates from 0.07 to 0.9g/L, protein from 0.15g/L to 0.9g/L, and lipids from 0.07 to 0.85g/L. Another study reported 297 per cent and 654 per cent increases in protein and carbohydrate concentrations after treating sewage sludge at 900W for 140 seconds (Yu et al, 2010). Under favourable conditions during microwave heating, especially those associated with high temperature (>180°C), caramelization and/or the Maillard reaction may occur, and these reactions can affect the odour and colour of treated HFS. ‘Caramelization’ is a non-enzymatic browning effect observed on short-chain polysaccharide sugars processed at elevated temperatures (Labuza and Baisier, 1992), while Maillard reactions are associated with reducing sugar’s reaction with amino acids known to generate dark brown-coloured complex compounds with characteristic smells (Penaud et al, 2000a; Dwyer et al, 2008). These were observed in a study where sewage sludge and HFS treated by microwave heating at 180°C and 200°C for 30 minutes generated a coal-like solid with a characteristic coffee-like and almond-like smell respectively (Afolabi et al, 2015).

Solubilizations of sulphur-containing compounds in HFS during microwave heating may also play a role in the reduction of odour (He et al, 2009). A study reported concentrations of dimethyl sulphide (8.1 ppm [v]) and methyl mercaptane (14.2 ppm [v]) in dewatered sewage sludge treated at 200°C falling to 11ppb (v) and 3ppb (v) in treated samples respectively (Namioka et al, 2011). In essence, reactive hydrolysis during microwave heating which renders sulphur-containing compounds non-volatile and dissolved into the liquid phase, coupled with a characteristic smell due to the thermal hydrolysis of carbohydrates and protein compounds and
subsequent intermediate reactions such as caramelization and/or the Maillard reaction, are likely mechanisms that can ensure odour eradication of HFS using microwave heating. Transformation of the foul odour associated with HFS could improve public acceptance of the microwave heating method.

4.2.2 Complete pathogenic deactivation

Microwave irradiation appears to be a viable option for generating environmentally safe materials that will satisfy existing WHO recommendations and other statutory regulations on pathogen deactivations. Disinfection studies on sewage sludge (as the closest alternative to real human faecal sludge in terms of pathogenic loads) at temperatures 70°C ≥ T ≤ 100°C and residence times (≤ 30 minutes) under microwave irradiation, have demonstrated efficacy, i.e. such treatment can effectively kill bacterial pathogens (notably \(E. \ coli\) and helminthic eggs) (Khalil and Villota, 1988; Hong et al, 2004; Martin et al, 2005; Pino-Jelcic et al, 2006). Furthermore, high-temperature treatments can deactivate xenobiotic and endocrine-disrupting compounds in biowastes (Wilson and Novak, 2009), producing end products that are free of biologically active organisms and bioactive exogenous compounds (Peterson et al, 2008). Microwave exposure for 1 minute at 65°C 1.34W h/g TS resulted in a sludge of coliform densities less than 1,000MPN/g total dried solids (Hong et al, 2006). Martin et al, (2005) also reported a similar observation at 85°C, where sewage sludge irradiated at 670W for 1 and 3 minutes resulted in complete destruction of coliforms to 33 and 2 total coliforms per 100ml respectively. At temperatures greater than 70°C, more studies have reported no microorganism activity, i.e. no detection of coliforms or \(E. \ coli\) in pre-treated sludge (Hong et al, 2004; Martin et al, 2005; Eskicioglu et al, 2009; Park et al, 2010).

Both the thermal and athermal effects of microwave heating at these temperature ranges play a significant role in pathogenic deactivation. The extents to which both effects contribute to pathogen kill are still relatively unknown. Selective heating, electroporation, cell membrane rupture and magnetic field coupling have all been reported as mechanisms behind the capacity of microwave heating to kill pathogens and enzymes (Kozempel et al, 1998). The selective heating and magnetic field coupling mechanisms suggest that the cell membranes of microbes are composed permeable lipids, which have a strong affinity for microwave absorption. Thus incident microwaves selectively energize such polar substances, causing irreversible damage to the cell membranes through DNA disruption. This subsequently leads to rupturing of the cell walls and membranes, and release of intracellular materials – leading to eventual denaturation
and death of the microbial cells (Belyaev et al, 1993; Kozempel et al, 1998). However, the electroporation and cell membrane rupture theories both seem to argue that electrical potential or voltage applied across the cell membrane causes a rupture or generation of pores, which subsequently results in leakage and release of intracellular materials (Kozempel et al, 1998). All the mechanisms point to the fact that microwave heating can attack coliform bacterial/microbial cell membranes, rupturing them and leading to their death. These theories have also been supported microscopically as SEM micrographs of raw and microwave irradiated sludgy material for 7 minutes (mins) at 2.45Ghz and 700W showed that – while raw sludge was highly porous and contained filamentous bacteria – those obtained from treated sludge showed structural rupture and disintegrated microbial cells (Ahn et al, 2009). Zhou et al, (2010) have also used SEM micrographs to confirm the disintegration of Gram-negative bacteria cell walls after microwave irradiation. Regardless of the mechanism(s) involved, it is evident that microwave heating can be highly effective and efficient in ensuring the destruction of pathogenic loads associated with HFS.

### 4.2.3 Enhanced dewaterability

Microwave heating can significantly improve the dewaterability of sludgy material such as HFS. Dewaterability essentially reduces sludge volume and is crucial for waste treatment technologies. About 40 per cent improvement in dewaterability was reported as a result of microwave heating of anaerobically digested sludge at 96°C (Eskicioglu et al, 2007b). Increasing the temperature to 175°C resulted in a 75 per cent improvement in dewaterability (Eskicioglu et al, 2008a). Similarly, significant reduction in capillary suction time (CST) – a measure of dewaterability – of up to 600 seconds has been reported from microwave-irradiated sludge treated at 175°C at 900W for a minute residence time (Toreci et al, 2009; Yu et al, 2009). To achieve sludge dewaterability, high microwave power (up to 900W), short residence times (less than 2mins) and temperature ranges over 120°C were recommended in another study (Yu et al, 2009).

Improved sludge dewaterability due to microwave heating has been attributed to both the thermal and athermal effects of microwave heating (Wojciechowska, 2005; Eskicioglu et al, 2007c, Toreci et al, 2009). Thermal effects result from direct coupling of electromagnetic energy with water molecules, which raise the temperature of sludgy materials, decrease their viscosity and enhance their filterability. In addition, microwaves can selectively energize chemically bound water in sludge to facilitate its chemical dissociation and release of bound water (Li and Noike, 1992). In effect, the molecular heating effect of microwave irradiation facilitates the rapid
disruption and disintegration of sludge flocs and bound water, and is responsible for improving
dewaterability within a shorter (less than 2mins in most cases) heating time (Wojciechowska,
2005; Yu et al, 2009). Athermal effects are attributed to the vibrational effects of microwaves on
hydrogen bonds in the cell walls of sludgy materials through the alternation of the electric field
of water (polar substance), causing overall weakening. This may also facilitate the breaking of
chemically bound water in sludgy materials – i.e. the continuous aligning of polarized parts of
materials’ macromolecules and water molecules with the poles of the electromagnetic field at
2.45 billion times per second, for example, initiates vibrational effects that could lead to
weakening and possible breakage of hydrogen bonds (Loupy, 2002; Eskicioglu et al, 2007c).
Understanding of the contribution/extent to which both affect dewaterability is still evolving,
but confirms microwave heating could enhance dewaterability of HFS.

4.2.4 Heavy metals recovery

Extraction and recovery of heavy metals can be both economical and environmentally attractive.
Detoxification of processed HFS of heavy metals promotes environmental compatibility and
ultimate prevention of soil and water pollution/contamination. Their recovery also makes them
available for reuse.

Microwave treatments are generally used in the presence of suitable acids to digest and enhance
the rapid dissolution of environmental samples for analysis/recovery, i.e. the microwave
digestion process facilitates the extraction and recovery of heavy metals. A study recorded up to
104 per cent metals recovery efficiency for Cd, chromium (Cr), Cu, Ni, Pb and Zn from sewage
sludge treated at 900W for a minute contact time (Jamali et al, 2009). Another also reported that
Cu ions were significantly leached from 179.4mg/l to 6.5mg/l in sewage sludge pre-treated at
600W for 3mins (Chen et al, 2005). Similar results were obtained with activated sewage sludge
pre-treated at 70°C for 90mins for the recovery of arsenic (As), molybdenum (Mb) (Danesh et al,
2008). Under favourable conditions, the thermal hydrolysis and subsequent solubilization
reactions associated with microwave processing of HFS, for example, could facilitate the
leaching of heavy metals, presenting them in ionic forms and effectively reducing the mobility of
metal ions in the aqueous phase. Full heavy metal recovery transcends HFS microwave
processing and may require additional processes, as reported in many studies (Perez-Cid et al,
1999; Perez-Cid et al, 2001; Kuo et al, 2005; Hsieh et al, 2008; Jamali et al, 2009). However,
microwaving HFS is potentially feasible for heavy metal digestion for extraction and recovery.
4.2.5 Nutrient recovery

Macromolecular protein and other nitrogenous compounds are sources of N in HFS. Microwave treatment can break down and enhance the solubilization of the protein components and other nitrogenous materials of HFS, i.e. microwaving HFS can flush N content in raw HFS into the liquor phase (as ammonia liquor concentrate) – supporting the proposition that human faecal sludge may be used as liquid fertilizer. At temperatures greater than 160°C, these compounds are hydrolyzed and decomposed to amino acids, organic-N and ammonium compounds. With increasing temperature (≥ 180°C), deamination and hydrolysis of amino acids into short-chain volatile fatty acids, ammonia and carbon IV oxide occurs (Sun et al, 2013). Fourier transform infrared spectroscopy (FTIR) and elemental CHN studies of thermally pretreated sewage sludge have further buttressed N delpetition into the liquor phase. FTIR analyses of thermally processed sewage indicated loss of N-bands seen in raw sewage into the liquor phase of processed sewage sludge (Grube et al, 2006; Smidt and Meissl, 2007), while CHN analysis suggests significant partitioning into the liquor phase of the original N percentage characterized in raw samples (Bagreev et al, 2001; Afolabi et al, 2015). Ammonia recovery may be seen as an apparent additional economic benefit from microwaving HFS. However, its direct use in agriculture requires further assessment as high concentrations of ammonia are environmentally damaging for rivers and watercourses, and use and dispersal of such concentrates require management and planning.

Ammonia recovered could also be combined with phosphates and could potentially be used to produce other solid plant fertilizers such as struvite (magnesium ammonium phosphates) for agricultural purposes. Phosphorus recovery is essential to achieving this. Generally, phosphorus tends to have an affinity for the solid phases. Only 10–15 per cent of P can be recovered from sewage sludge treated at 180°C–240°C for up to 90mins (Sun et al, 2013). In addition to temperature conditions >170°C, catalysts are required to ensure P solubilization and release as phosphates of calcium and iron into the liquor phase prior to subsequent crystallization/precipitation into fertilizers such as struvite. For example, a study used hydrogen peroxide as a catalyst with microwave irradiation at 171°C for 5mins and was able to recover up to 84 per cent of P from sewage sludge (Liao et al, 2005). Similar results were obtained in another study where acids hydrolysis in the presence of hydrogen peroxide was used to recover 61 per cent of total P and 36 per cent of total Kjeldahl nitrogen (TKN) at 120°C for 5mins (Wong et al, 2006). Although, microwaving HFS provides an opportunity for P recovery,
additional downstream processing maybe required for higher recovery rates of phosphates suitable for agricultural use.

Comparable to N content, microwaving HFS at temperatures greater than 180°C can also favour the solubilization of K into the liquid phase. Up to 70 per cent K was recovered in liquid products from sewage sludge hydrothermally treated between 180°C and 240°C; this reported a preferential partitioning of N and K into the liquor phase, in contrast with P which has a high affinity for the solid phase (Liao et al, 2005; Sun et al, 2013). Nonetheless, the retention of P in solid product chars may be seen as an advantage, for P is a nutrient for crop growth. If used as solid fuel, phosphorus can be recovered in the ash content of combusted chars (Escala et al, 2012). Consistent with industrial practices, the prices of 1kg N (as TN), 1kg P (as K₂O) and 1kg K (as KCl) in commercial fertilizers has been up to US$13.22, US$11.27 and US$6.63 respectively over the last decade.² Hence the recovery of nutrients from HFS represents a feasible financial benefit to the whole faecal sludge management chain.

4.2.6 Carbon and energy recovery

Carbon recovery is essential for diverse uses including energy, sequestration and fixation into agricultural soil. Under favourable conditions such as those associated with the carbonization process – i.e. heating at temperatures 160°C–250°C in the absence of oxygen under autogenous pressure (Peterson et al, 2008; Libra et al, 2011) – microwave heating can be used to process HFS to recover energy-densified solids called chars (Wahi et al, 2006; Tian et al, 2013; Afolabi et al, 2015). These chars are characterized by a significantly higher heating value of up to 25MJ/kg (as against 19MJ/kg recorded in raw HFS) (Afolabi et al, 2015); and are coal-like in appearance and highly porous (which enhances air distribution during combustion), making them suitable candidates for fuels. The relatively higher heating value (HHV) and energy yield from chars unlocks opportunities for co-combustion with low-rank coals (with similar ranges of HHV) (Speight, 2005) for energy generation, although thermal compatibility requires further and careful assessment (Parshetti et al, 2013). Even at higher temperatures, microwave pyrolysis of sewage sludge has yielded other biofuels such as bio-oil and gaseous hydrocarbon (Menendez et al, 2002; Tian et al, 2013). Wahi et al, (2006) in their study, microwaved sewage sludge at 650°C and recovered 28 per cent chars, 6 per cent bio-oil and 68 per cent gaseous hydrocarbon, with HHV of the bio-oil and chars being 28.9MJ/kg and 12.4MJ/kg respectively. These are similar results to

² Values were obtained from studying price variations for commodity fertilizers over ten years at: http://www.indexmundi.com/commodites/
those of many other studies that have demonstrated the efficacy of using the microwave process to generate fuel in the forms of chars, bio-oil and gaseous hydrocarbons from sewage sludge at temperatures ranging between 200°C and 1000°C (Dominguez et al 2003, 2005, 2006; and 2008).

Chemical oxygen demand (COD) indicates the fractions of carbon transferred into the liquid phase from hydrolysis and solubilization of organic and inorganic components during thermal treatment, such as those associated with microwave irradiation. Apart from disposal processes, soluble COD is a helpful measure for assessing liquor use as a substrate for anaerobic digestion (AD) to recover biogas fuel (Bougrier et al, 2008; Tyagi and Lo, 2011). Microwave has been studied and used extensively to increase the concentration of soluble COD in sludgy materials especially for AD (Zheng et al, 2009; Eskicioglu et al, 2006; Eskicioglu et al, 2007a; Yeneneh et al, 2015). COD obtained for microwave analyses of HFS liquors ranges between 21g/L to 26g/L (Afolabi et al, 2015) and is comparable with those recovered from sewage sludge studies (Dwyer et al, 2008; Wilson and Novak, 2009; Eskicioglu et al, 2009), indicating utility for anaerobic digestion. In general, temperatures in the range 160°C to 180°C are considered to be optimal for generating AD feedstock from biowastes, based on COD concentration (Li and Nioke, 1992; Wilson and Novak, 2009). Higher temperatures were reported to decrease the biodegradability of sludge (Stuckey et al, 1978; Bougrier et al, 2008).

Future research in this area could involve the standardization of optimum temperature range, microwave power and residence time required to achieve both HFS treatment and value recovery. This will be crucial for the design of microwave-based sanitation facilities.

4.3 Comparing microwave irradiation with conventional heating for processing HFS

Microwave heating potentially represents a better heating approach for processing HFS over conventional conduction heating methods. Specifically, microwave heating causes selective and rapid volumetric heating and faster reaction rates, which make it superior to conventional heating (Clark et al, 2000; Dominguez et al, 2006; Yin, 2012). The main differences between microwave and conventional heating processes are summarized in Table 2. The overarching implications of these can be seen in time and energy savings and associated costs of processing.

The conventional heating process is characterized by an external supply of heat transferred by thermal gradients via conduction, convection and radiation to heated material, i.e. the heat supplied penetrates from the surface towards the core of the heated material, causing the outer
region of heated components (i.e. reactor vessels and material surfaces) to be hotter than their inner regions. This is inefficient when compared with what occurs during microwave heating. When microwaves heat materials, selective heating of dipoles occurs throughout the heated material (as microwaves can penetrate and deposit energy throughout the volume of the material), causing a uniform and rapid volumetric heating effect. As a result, the inner regions of the heated material become hotter and this reduces the chance of surface drying/degradation, which is commonly associated with conduction heating (Remya and Lin, 2011). In addition, since most microwave handling components are good insulators, e.g. quartz or glass, this causes a reverse thermal gradient – i.e. a higher temperature at the core and lower temperature at the surface (Clark et al, 2000). Also, in conventional processes, heating is slower because (source energy such as electric) energy is first converted into heat energy, before being transferred to the sample material from the conducting surface. In microwave heating, microwave energy is delivered directly to the heated material through molecular interactions with the electromagnetic field. Heat is induced within the core of the material through the direct conversion (rather than transfer of heat) of microwaves into calorific heat, thereby turning the material into a heat source. As a result, microwave heating provides faster and more energy-efficient heating. This feature has been linked with the accelerated reaction rates microwave heating offers, because turning the heated material to a heat source can lead to an alteration of reaction kinetics with energy-saving implications (Appleton et al, 2005; Sobhy and Chaouki, 2010). Conventional processes also tend to require the construction of processing equipment in situ. Using microwave heating can overcome this problem due to the relatively lower footprint of the technology. Reactor vessels used for conventional processes are usually thick-walled metallic conductors, which must withstand pressurized conditions and sometimes the corroding environment associated with high-temperature processes. This may have cost implications, especially when scaling the process. However, with microwave heating this can be avoided, as metals are poor microwave conductors; hence cheap and readily available insulators such as ceramic and other refractory materials can be used to design treatment/processing reactor vessels.
Table 2 Microwave versus conventional heating

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Microwave heating</th>
<th>Conventional heating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy</td>
<td>Conversion of energy from electromagnetic to heating, i.e. energy transfer</td>
<td>Transfer of heat associated with thermal gradients, i.e. heat transfer</td>
</tr>
</tbody>
</table>
| Heating                 | 1. In-core volumetric and uniform heating, simultaneously heating materials due to energetic coupling at molecular level  
                            2. Non-contact heating                                                            | 1. Superficial heating via conduction / convection                                   
                            2. Need for contact between heating source and heated material                    |
| Rate                    | Rapid and efficient rate of heating due to molecular interaction at a very high frequency; hence desired peak temperature can be reached quicker | Slow, inefficient, limited by material thermal conductivity                            |
| Selectivity             | Highly selective for polar substances, but poor for non-polar materials, and hence can trigger new products | Non-selective                                                                        |
| Hotspots                | Localized hotspots due to in-homogeneities of microwave electric field concentration or due to dielectric properties within material; this causes arcing, i.e. temperature within material to be higher than bulk temperature of material | No hotspots                                                                          |
| Dependence on nature of heated materials | Highly dependent on dielectric properties of the material  | Less dependant                                                                        |
| Heating control features | Enhanced control and process monitoring features; precise on/off power and temperature controls; energy input can be stopped abruptly | Poor controllable nature                                                              |
| Cooling rate            | Faster cooling rates due to enhanced cooling units                                 | Longer cooling rates                                                                  |
| Particle size           | Not sensitive to particle size                                                     | Highly sensitive to particle size in terms of enhancing heating rate                  |

5. Key technological and design parameters considerations for microwave technology crucial for scaling microwave for sanitation purposes

Most of the successes and feasibility studies conducted in the use of microwave irradiation for processing biomass/biowastes are largely laboratory based, involving relatively small quantities of samples. As with any technology, there are parameters that must be duly considered for optimization and design at various stages of scaling. These parameters are considered below.
5.1 Dielectric properties of HFS components

Microwave heating induces physicochemical and structural transformation as materials are being processed, and these transformations affect dielectric properties – and by extension the ability of microwaves to generate heat varies during the process. It is necessary to determine the dielectric properties (dielectric capacity and loss) of human faeces or urine over microwave frequency ranges and at different temperature regimes, particularly those associated with HFS treatment and product recovery to optimize the microwave heating efficiency. Determination of the dielectric properties depends on many factors such as temperature changes, microwave frequency, moisture content, the chemical composition of material, density of material and physical characteristics of material (homogeneity and particle size distribution) (Thostenson and Chou, 1999), and these may present a significant developmental challenge for highly heterogenous and variable HFS. Furthermore, key parameters such as power absorption (i.e. the amount of transmitted microwave power absorbed per unit volume by heated materials from the electromagnetic microwave distribution) and heating rate (i.e. measures of the amount of power absorbed that is converted to heat within the material) – which are crucial for enhancing microwave-heating efficiency – are largely dependent on the dielectric properties of materials irradiated, as shown in Equations 2 and 3 (Venkatesh and Raghavan, 2004; Kappe, 2004).

\[ P_v = 2\pi f \varepsilon_0 \varepsilon' \tan(\delta) |E|^2 \]  
Equation 2

Where  
- \( P_v \): Power absorbed per unit volume of material, W/m³  
- \( f \): Frequency of microwave, Hz  
- \( \varepsilon_0 \): Permittivity of free space = 8.85 x 10⁻¹² F/m  
- \( \varepsilon' \): Dielectric constant  
- \( \tan(\delta) \): Tangent loss, where \( (\delta) \) is the loss angle  
- \( E \): Electric field, V/m

\[ \frac{\Delta T}{\Delta t} = \frac{P_v}{\rho C_p} \]  
Equation 3

Where  
- \( \Delta T \): Change in temperature, K  
- \( \Delta t \): Residence time involved, Seconds  
- \( \rho \): Density of the material, kg/m³  
- \( C_p \): Heat capacity of the material, J/K  
- \( P_v \): Power absorption of material, W/m³
5.2 Penetration depth

Penetration depth, $D_p$, is the distance at which transmitted microwave power drops to $\exp^{-0.368}$ (equivalent to 36.8 per cent) from its original value at the surface of the heated material. Beyond this depth, volumetric heating due to microwave energy is insignificant. This parameter is particularly important for determining the uniformity of heating in a material. Penetration depth is estimated as:

$$D_p = \frac{3\lambda_0}{8.686\pi\tan\delta \left(\frac{\varepsilon'}{\varepsilon_0}\right)}$$

Equation 4

and is related to frequency of microwave irradiation as:

$$D_p = \frac{c\varepsilon_0}{2\pi\varepsilon e}$$

Equation 5

Where $\lambda_0$ and $c$: Incident wavelength of microwave (m) and speed of light (m/s)

It follows that penetration depth is dependent on the frequency of microwaves and the dielectric properties of an irradiated material. Higher frequencies and larger dielectric properties will produce low penetration depth and cause on-surface heating of the material, while lower frequencies and smaller dielectric properties of a material will promote volumetric heating (Clark et al, 2000; Tyagi and Lo, 2013a). For instance, penetration depths for 915MHz microwaves are about three times larger than for 2.45GHz microwaves. This is the main reason for the wider application of this frequency in large-scale industrial systems.

In addition, penetration depth is inversely proportional to absorption of microwaves in a material. This means that the size of treated material, for instance the size of the container holding waste, should be about the same as the penetration depth. It follows that if the penetration depth of incident microwaves is less than the thickness of the sample, only the surface of the material will be irradiated and heated, while the rest will be heated by conduction (Yin, 2012). In contrast, if the size of the sample is much smaller than the penetration depth, most of the microwaves will go through the sample without being absorbed.

Table 3 further relates the temperature and moisture content dependence of penetration depths. From Table 3, cold water absorbs microwaves about four times stronger than hot water. This means that colder parts will warm up faster, resulting in a more uniform heating. Table 3 also suggests that increasing moisture content and temperature may increase penetration depth. More
studies are needed on optimized penetration depth for both moisture content level and microwave temperature of treatment.

Table 3 Microwave energy penetration depth at 2.45GHz (extracted from Decareau 1985; Puschner Microwave Power Systems, 2016)

<table>
<thead>
<tr>
<th>Materials</th>
<th>Penetration depth (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material with high moisture (≥50%)</td>
<td>3.1</td>
</tr>
<tr>
<td>Material with intermediate moisture</td>
<td>4.4</td>
</tr>
<tr>
<td>Material with low moisture (≤30%)</td>
<td>8.2</td>
</tr>
<tr>
<td>Water at 25°C</td>
<td>1.4</td>
</tr>
<tr>
<td>Water at 95°C</td>
<td>5.7</td>
</tr>
<tr>
<td>Ice at -12°C</td>
<td>1,100</td>
</tr>
<tr>
<td>Sewage sludge (96% moisture)</td>
<td>0.2*</td>
</tr>
<tr>
<td>HFS (95% moisture)</td>
<td>1.9*</td>
</tr>
</tbody>
</table>

*Unpublished experimental data determined at 15+/−2°C

5.3 Cold and hot spots, effective electromagnetic coupling and microwave power efficiency

As microwave energy is transferred by the electromagnetic field, non-uniformity within the electromagnetic field may result in non-uniform heating (Thostenson and Chou, 1999; Remya and Lin, 2011). For example, microwave ovens use standing waves confined inside a metal resonant cavity. This means that microwave amplitude and power generation density are functions of position. Cavity locations where amplitudes of standing waves are near maximum will experience much larger heat generation density as compared to locations where microwave amplitude is near zero. This effect causes hot and cold spots during microwave irradiation.

The existence of hot and cold spots may affect the design of the toilet waste treatment system, if the cavity is not properly designed to alternate and expose material to uniform irradiation. Batch systems or semi-continuous systems, where the waste moves in discrete batches, may also encourage hot/cold spots generation. Waste in cold spots may experience incomplete pathogen deactivation, especially if mixing of waste is not sufficient to average out microwave exposure. A common approach to eliminate hot and cold spots is to use a turntable, as obtained in kitchen microwaves. The turntable moves irradiated materials between hot and cold spots and provides an averaged and more uniform heating. An alternative approach, although not frequently used, is to employ a rotating metal reflector or stirrer installed inside of the microwave cavity. The reflector/stirrer effectively shifts the location of hot/cold spots, allowing for uniform heating.
One of the challenges of microwave heating is to ensure efficient absorption, i.e. coupling between microwaves and the treated material. Understanding the generation, propagation and interaction of microwaves with materials is critical to ensuring effective coupling. Reflection of microwaves at material interfaces, too short or too long microwave penetration depth, and/or overall poor design of the microwave cavity may result in poor coupling. This requires careful assessment, to prevent significant impact on the advantageous kinetics of microwave HFS processing. Even with perfect coupling, a microwave magnetron will convert only about 50 per cent (for 2.45GHz) and 85 per cent (for 915Mhz) of the electrical energy into microwaves (Engelhart, 1992; Haque, 1999), while the remainder is dissipated as heat inside of the magnetron source. This conversion efficiency is apparently dependent of microwave frequency and should be considered when designing off-grid microwave-based sanitation systems for HFS processing.

5.4 Temperature and pressure measurements

The challenge of precise temperature measurement within the core of the heated material during the microwave process has also been highlighted in a few reports (Tian et al, 2013; Hoseinzadeh-Hesas et al, 2013). Most measurements are conducted externally via infrared pyrometers. Effective temperature monitoring within the core of heated HFS and autogenous pressure build-up from the exothermic reactions associated with high temperature treatment of HFS, is crucial for modelling and control designs. Other important factors – such as cavity design, choice of materials to design reactors for processing HFS, and process conditions that will achieve both treatment and value recovery – also require careful assessment during design and scaling of microwave technology for HFS processing.

Conclusion

This review discussed the characteristic value compositions of HFS as a significant nutrient-based resource and explored microwave heating as viable method that can both treat and recover value from HFS. The high-moisture content (up to 97 per cent) that characterizes HFS makes it fit the dielectric heating spectrum, and may trigger very rapid and volumetric heating, promote novel reaction pathways and enhance reaction rates. Enhanced reaction rates present an opportunity to develop higher throughputs. Microwaves can potentially eradicate foul odours and kill pathogens efficiently, thereby making products safe to handle. This may further boost public acceptability of the technology.
The highly selective nature of microwave heating targeting dipoles can trigger novel reactions that can lead to molecular and structural changes in processed HFS, and thus provide for the generation of new products. Ammonia liquor concentrate, and biofuel such as chars, bio-oil and biogas products for diverse applications in agriculture and energy generation, are examples of the valuable products that are potentially recoverable from microwaving HFS. Successful dewatering, plant nutrients and fuels recovery from microwaving HFS could improve the financial incentives for waste management in both developed and developing countries, if the contingent sustainability value of the process is assessed.

However, crucial technological factors – such as the dielectric properties of HFS components (especially at high temperatures, which may be required to achieve both treatment and value recovery) and the physics of the microwave/HFS interactions during heating – are necessary investigations to optimize the efficiency of the microwave heating process. As the microwave processing equipment will determine the electromagnetic field, the design of microwave equipment (especially the cavity design) is also particularly important. Hence, more work on system designs, development and performance improvement of the microwave technology, outside laboratory applications, is necessary to feed into the integration of the technology for sanitation purposes.

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