EXPERIMENTAL STUDY AND NUMERICAL MODELLING OF SELF-HEATING BEHAVIOUR OF TORREFIED AND NON-TORREFIED BIOMASS FUELS

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Abstract

Biomass pre-treatment using torrefaction technology has been proven to upgrade energy density, thermal and physical properties of the biomass. The torrefied biomass fuel has higher energy content, which could be comparable to low-rank coal. However, similar to coal, self-heating is one of the major safety concerns during the storage and handling, which can lead to ignition. There are several events of accidental fires that occurred in the past, resulting from self-heating leading to ignition. This work focuses on the experimental study and numerical modelling of self-heating behaviour of microwave torrefied biomass and non-torrefied biomass stored in a large pile. Physical and chemical properties of the materials were determined using proximate and ultimate analysis. The calorific values of the non-torrefied sample are 18.4MJ/kg and 22.0MJ/kg for torrefied biomass. Self-heating propensity is determined based on heat evolution and mass changes analysis using the thermogravimetric analysis to determine the thermal decomposition behaviour and thermal decomposition kinetics parameters of the torrefied and non-torrefied material. The activation energy for thermal decomposition in the air is in between 64.5 to 84.4 kJ/mol for the non-torrefied sample and 53.1 to 70.7 kJ/mol for the torrefied sample. However, the activation energy for thermal decomposition in nitrogen is between 58.7 to 70.9 kJ/mol for non-torrefied sample and 59.2 to 71.8 kJ/mol for the torrefied sample. The evaluation of the self-ignition risk was done based on the characteristic oxidation temperature and the activation energy. The activation energy for torrefied biomass fuel indicated that the material is reactive and have a higher tendency to self-heating. The risk ranking graph; is drawn based on the activation energy and characteristic temperature of the samples against other materials from literature. Both samples are classified as a medium risk, but with microwave torrefied sample in the rank of higher tendency to self-ignite comparing to the non-torrefied sample. A series of bulk test was performed to investigate the heating behaviour on a larger scale to examine the effect of the bulk size and oven temperature. The test showed that when the oven temperature is below and at 180°C no ignition is detected for the non-torrefied sample. However, microwave torrefied sample started to self-heat at 4002 seconds (66.7 minutes). It can be concluded that the torrefied sample is more reactive compared to the non-torrefied sample. Series of
the bulk test is carried out in the uniformly heated oven at constant air temperature to study the thermal heating behaviour independently. The test showed the ignition induction time for microwave torrefied sample heated at 180°C is 4002 seconds at Bulk 1, 5066 for Bulk 2 and 7611 for Bulk 3. The ignition induction time increased with the decreasing of the bulk volume. The ignition induction time is decreasing when the oven temperature is increased. A simulation model to predict the heating behaviour of the materials, in an open storage area, had been developed using COMSOL Multiphysics® software. The numerical model was used to examine the thermal behaviour of the pile based on coupled heat and mass transfer in porous media, which includes kinetic parameters, obtained using thermogravimetric analysis. The numerical model was validated against the results of the bulk tests with good agreement. Simulations were carried out to examine the effect of the height and ambient temperature on the thermal behaviour of the pile. The simulations demonstrated that the ignition induction time decreased when the ambient temperature increased. The ambient temperature of 60°C is established as critical ambient temperature for the storage of microwave torrefied sample, while 80°C is the critical ambient temperature for non-torrefied sample. Based on the simulation, the biomass needs to be piled up vertically instead of horizontally to avoid the multiple hot spots. The results from this study can be used as decisional support information towards achieving the safe storage of torrefied biomass.
Acknowledgement

I would like to express my most sincere gratitude and appreciation to Dr Yajue Wu, who has patiently mentored me throughout my PhD journey. She was supportive and helpful during the stressful and toughest time in my life. She has given me insightful guidance as well as encouragement for the last four years.

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To my family - thank you for the unconditional support and believing in me:
~ Ayah, Mama, Abang, Kak Huda, Azmin, Elena, Firdaus, Munirah, and Faiz ~

Lastly, thank you to all my amazing friends who always by my side...

“Success isn’t a result of spontaneous combustion. You must set yourself on fire.”

– Arnold H. Glasgow
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## Abbreviations

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<th>Abbreviation</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>SIT</td>
<td>Self ignition temperature</td>
</tr>
<tr>
<td>TMWL</td>
<td>The maximum weight loss</td>
</tr>
<tr>
<td>DTA</td>
<td>Differential thermal analysis</td>
</tr>
<tr>
<td>DSC</td>
<td>Differential scanning calorimeter</td>
</tr>
<tr>
<td>TGA</td>
<td>Thermogravimetric analysis</td>
</tr>
<tr>
<td>TG</td>
<td>Thermogravimetric</td>
</tr>
<tr>
<td>DTG</td>
<td>Differential thermogravimetric</td>
</tr>
<tr>
<td>CP</td>
<td>Crossing point</td>
</tr>
<tr>
<td>TIC</td>
<td>Temperature of Initial Combustion</td>
</tr>
</tbody>
</table>
## Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Pre-exponential factor</td>
<td></td>
</tr>
<tr>
<td>Bi</td>
<td>Biot number</td>
<td></td>
</tr>
<tr>
<td>$C_p$</td>
<td>Specific heat</td>
<td>J/(kg K)</td>
</tr>
<tr>
<td>$D_p$</td>
<td>Particle diameter</td>
<td>mm</td>
</tr>
<tr>
<td>$E_a$</td>
<td>Activation energy</td>
<td>J/mol</td>
</tr>
<tr>
<td>H</td>
<td>Height</td>
<td>m</td>
</tr>
<tr>
<td>$H$</td>
<td>Convective heat transfer coefficient</td>
<td>W/(m² K)</td>
</tr>
<tr>
<td>$K$</td>
<td>Reaction rate constant</td>
<td>min⁻¹</td>
</tr>
<tr>
<td>$L$</td>
<td>Length</td>
<td>m</td>
</tr>
<tr>
<td>Q</td>
<td>Heat source</td>
<td>W/m³</td>
</tr>
<tr>
<td>R</td>
<td>Universal gas constant, 8.314 J mol⁻¹ K⁻¹</td>
<td></td>
</tr>
<tr>
<td>$q^\prime$</td>
<td>Heat source</td>
<td>J/kg</td>
</tr>
<tr>
<td>$T$</td>
<td>Temperature</td>
<td>K</td>
</tr>
<tr>
<td>$t$</td>
<td>Time</td>
<td>s</td>
</tr>
<tr>
<td>V</td>
<td>Volume</td>
<td>m³</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Heating rate</td>
<td>K/min</td>
</tr>
<tr>
<td>$w$</td>
<td>Weight original sample</td>
<td>mg</td>
</tr>
<tr>
<td>$w_o$</td>
<td>Weight at the start of the stage</td>
<td>mg</td>
</tr>
<tr>
<td>$w_\infty$</td>
<td>Weight at the end of the stage</td>
<td>mg</td>
</tr>
<tr>
<td>V</td>
<td>Volume</td>
<td>m³</td>
</tr>
<tr>
<td>$O_2$</td>
<td>Oxygen concentration</td>
<td>mol m⁻³</td>
</tr>
</tbody>
</table>

### Greek symbols

- $A$: Thermal diffusivity, m²/s
- $\lambda$: Thermal conductivity, W/(m K)
- $P$: Bulk density, Kg/m³
- $\varepsilon$: Porosity
- $\delta_c$: Damkohler number.

### Subscript

- $F$: fuel
- $\text{react}$: reaction
- $\text{conv}$: convective
- $\text{crit}$: critical
- $\text{eff}$: effective
- $\text{charac}$: characteristic
Chapter 1: Introduction

1.1 Background

Biomass is a source of renewable energy that has a huge potential to play a major role in the energy sector as alternative fuels to coal. Biomass is defined as “the biologically degradable fraction of products, waste and residues from the biological origin of agriculture that include vegetable and animal substances, forestry and related industries including fisheries and aquaculture, as well as the biodegradable fraction of industrial and municipal waste” (Renewable Energy Directive, 2009). However, despite many sources of biomass, lignocellulosic biomass has been considered as the most promising energy source due to its surplus availability, usage practicality, and relatively low cost. The lignocellulosic biomass refers to plant biomass that is composed of cellulose, hemicellulose, and lignin.

![Figure 1.1: The carbon neutral cycle (Vassilev et al., 2015).](image)

In addition to that, lignocellulosic biomass is the only carbon neutral energy source that does not contribute to the greenhouse effect. Therefore lignocellulosic biomass has become the most favourable source due to its carbon neutral advantage. The neutral carbon cycle of carbon dioxide is shown in Figure 1.1, where the neutralisation of carbon dioxide emission achieved when converting biomass to
energy since the growth of new biomass removes carbon dioxide from the atmosphere. Unless otherwise stated, biomass is referred as lignocellulosic biomass throughout this thesis.

As mentioned earlier, the main components of lignocellulosic biomass are cellulose, hemicellulose, lignin and extracts. Thus, cellulose is a component with a crystalline structure that is insoluble in water as well as resistant towards depolymerisation (Badiei et al., 2014). Hemicellulose is a backbone of the plant cell wall that has sugar units such as xylose, mannose, galactose, arabinose as well as glucose (Eriksson, 2011). Lignin provides mechanical strength to the plant cell wall by covalent linkage. Lastly, terpene, phenol and different types of fats are among extractives in the lignocellulosic biomass component. Different types of biomass will have a different component breakdown. The examples of biomass components adopted from study by Mašek et al. (2013a) are shown in Figure 1.2.

![Figure 1.2: Components in (a) willow chips (b) straw pellets (Mašek et al., 2013a)](image)

The biomass characteristics are influenced by its origins from which it has been collected. Biomass may have the same major component but in different proportion, which results in a broad range of fuel properties. According to Naik et al. (2010), the same type of biomass can have different composition based on the climatic condition as well as the seasonal variation. Badiei et al. (2014) added that the composition of biomass component is also affected by the types of the biomass such as hardwood or softwood. The work by Vassilev et al. (2010) recognised various factors that affected the composition of biomass such as the growth conditions, a distance of the plant from the source of pollutions, harvesting time, types of biomass including plant
species or part of the plants. Its chemical composition cannot be defined precisely for a given tree species or even for a given tree properties. The utilisation of each biomass is also different from one to another. Table 1.1 shows the examples of biomass subgroup, species and varieties coniferous as well as their example of usage. It can be witnessed from the table that biomass is a versatile fuel, which is very different from one to another in term of utilisation.

Table 1.1: Lignocellulosic biomass groups and examples of utilisation (Vassilev et al., 2010; AEBIOM, 2012)

<table>
<thead>
<tr>
<th>Groups</th>
<th>Biomass sub-groups, species and varieties coniferous</th>
<th>Example of utilisation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wood and woody biomass</td>
<td>Coniferous or deciduous, angiospermous or gymnospermous and soft or hard such as stems, branches, branches (twigs), leaves (foliage), bushes (shrubs), chips, lumps, pellets, briquettes, sawdust, sawmill and others from various wood species</td>
<td>Wood chips &amp; particles, firewood briquettes and pellets for energy source</td>
</tr>
<tr>
<td>Herbaceous and agricultural biomass</td>
<td>Annual or perennial, arable or non-arable and field-based or processed-based biomass from various species such as grasses and flowers, straws, stalks, fibres, Shells and husks, pit and other residues from numerous sources</td>
<td>Oilseed crops: for biodiesel production. Sugar &amp; starch crops: for bioethanol. Lignocellulosic &amp; woody crops: for heat and power production; second generation biofuels</td>
</tr>
<tr>
<td>Aquatic biomass</td>
<td>Marine or freshwater, macroalgae or microalgae and multicellular or unicellular species</td>
<td>Algae: production of biogas for energy production</td>
</tr>
</tbody>
</table>

Fossil fuels such as coal and oil have been dominated the world energy supplies for decades. Based on International Energy Agency (IEA) data from the World Energy Balances (2016) presented in Figure 1.3, the global primary energy supply had increased by almost 2.5 times from 1971 to 2014. The oil supply decreased to 31% from 44% in 1971, while coal supply increased 3% in 2014. The supply of natural gas also increased from 16% in 1971 to 21% in 2014. The trend shows that world
primary energy supplies by some means still depend on fossil fuel with 81% of energy supply in 2014 is from fossil fuels.

![Figure 1.3: World total primary energy supply based on the types of fuel. (IEA, 2016)](image)

Unfortunately, harmful emission such as SO₂, NOₓ, CO₂, N₂O, particulate matter, mercury, cadmium and other acid gases are being released into the environment by the coal-fired power plant (Nalbandian, 2010). The burning of fossil fuels such as coal that cause the harmful emission started since the Industrial Revolution as early as the 1800s has been recognised as one of the main contributors to the increase of the greenhouse gases concentration in the atmosphere especially carbon dioxide (Life Science, 2016).

Coal is a sedimentary rock that composed of lithified plant materials remains that consist of carbohydrates, lignin, proteins and other polymers including hemicelluloses, suberin as well as cutin (Speight, 1994). Coal can be grouped into types based on its rank, which is a measure according to the degree of organic metamorphism (changes). The rank of coal is from lowest to highest are peat, lignite, sub-bituminous, bituminous and anthracite.

The chemical composition of coal is different from biomass, especially the amount of hydrogen, oxygen and carbon. In comparison to biomass, coal has much higher carbon content and fewer fractions of hydrogen and oxygen. Thus, coal is a fuel with a low atomic ratio of oxygen to carbon (O/C); which is more favourable in energy conversion. The low O/C fuel is favourable because of its higher gasification
efficiencies (Prins et al., 2006b) and decrease amount of smoke and water vapour (Kambo & Dutta, 2014). Furthermore, biomass has higher volatile matters compared to coal, which could lead to low ignition temperature causing more amounts of combustible gas production during thermal conversion process (Chiang et al., 2012).

Based on Table 1.2, the energy density of coal is considerably higher than the one in wood as well as solid biofuel such wood pellet and torrefied pellet. As mentioned earlier, biomass fuels have more oxygen and hydrogen and less carbon than coal. Thus, the properties are agreeable with the fact that biomass fuels have much higher volatile matters and moisture content compared to coal as well as lower heating value. Based on Table 1.2, the biomass fuel that undergone the pre-treatment such as torrefaction had improvement in properties, for example the heating value of the torrefied pellets is higher compared to the wood pellets. The improved properties are comparable to coal. Thus, the positive outcome from the biomass pre-treatment is creating the demand for solid biofuels in the energy industry to fulfil the carbon neutral energy demand.

Nevertheless, apart from the climate change problem due to harmful emissions from the combustion of fossils fuels, the exhaustion of fossil fuels and price increases resulted in the necessity to shift to the renewable energy source such as biomass. In light of that, European Union (EU) had set a binding target for a renewable energy source to contribute 20% of the total energy needs by 2020 (Renewable Energy Directive, 2009). These developments have promoted the growth of renewable energy usage.

Figure 1.4: Source of total energy consumption in Europe in 2014 (AEBIOM, 2016)
Table 1.2: Properties of biomass fuels and coal (Nunes et al., 2014\textsuperscript{1}; Koppejan et al., 2012\textsuperscript{2})

<table>
<thead>
<tr>
<th></th>
<th>Wood\textsuperscript{2}</th>
<th>Wood pellets\textsuperscript{2}</th>
<th>Wood chips\textsuperscript{1}</th>
<th>Torrefied pellets\textsuperscript{2}</th>
<th>Bituminous Coal\textsuperscript{1}</th>
<th>Coal\textsuperscript{2}</th>
<th>Charcoal\textsuperscript{2}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (%)</td>
<td>30-45</td>
<td>70-10</td>
<td>30-60</td>
<td>1-5</td>
<td>5-10</td>
<td>10-15</td>
<td>1-5</td>
</tr>
<tr>
<td>Volatile matter (% db)</td>
<td>70-75</td>
<td>70-75</td>
<td>70-75</td>
<td>55-65</td>
<td>-</td>
<td>15-30</td>
<td>10-12</td>
</tr>
<tr>
<td>Lower heating value (MJ/kg)</td>
<td>9-12</td>
<td>15-18</td>
<td>6-13</td>
<td>20-24</td>
<td>&gt;25</td>
<td>23-28</td>
<td>30-32</td>
</tr>
<tr>
<td>Bulk density (kg/m\textsuperscript{3})</td>
<td>200-250</td>
<td>550-750</td>
<td>250-400</td>
<td>750-850</td>
<td>800-10000</td>
<td>800-850</td>
<td>~200</td>
</tr>
<tr>
<td>Energy density (GJ/m\textsuperscript{3})</td>
<td>2.0-3.0</td>
<td>7.5-10.4</td>
<td>2.5-3.2</td>
<td>15-18.7</td>
<td>20-25</td>
<td>18.4-23.8</td>
<td>6-6.4</td>
</tr>
<tr>
<td>Biological degradation</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Hygroscopic properties</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
<td>Hydrophobic</td>
<td>Hydrophobic</td>
<td>Hydrophobic</td>
<td>Hydrophobic</td>
</tr>
<tr>
<td>Dust</td>
<td>Average</td>
<td>Limited</td>
<td>Average</td>
<td>Limited</td>
<td>Limited</td>
<td>Limited</td>
<td>Limited</td>
</tr>
</tbody>
</table>
Based on Figure 1.4, 16% of the energy consumption originated from the renewable sources (biofuels and other renewable energy sources) in 2014. That current status is a good sign showing that the target set in the Renewable Energy Directive, (2009) is achievable. Therefore, a new target had been set by the European Union in the 2030 Energy Policy, where the final energy consumption in the European Union at least 27% by 2030 (Energy,2017).

In Figure 1.4, biofuels refer to liquid fuels produced from biomass such as biodiesel and bioethanol. While other renewable sources include the wind, hydro, solar, geothermal and bioenergy. Therefore, the energy produced from biomass sources is called bioenergy. Bioenergy is widely used in heating and cooling, power generation as well as transport application. Due to the high availability as well as the consistency of supply around the world, the biomass is an excellent material compared to other sources. The growth of biomass trade especially the trade of pellets and biofuels is moving towards a more globally traded commodity. In Europe, wood pellet consumption reached 20.3 million tonnes that represent 6% of total solid biomass usage (AEBIOM, 2016).

However, the direct use of lignocellulosic biomass is not possible for energy conversion due to several issues such as handling. Therefore, the lignocellulosic biomass needs to be transformed into fuel in solid, liquid or gas form using various pre-treatment technologies. Wood pellet is one of the solid biofuels that can be produced from the pre-treatment of lignocellulosic biomass. Wood pellet is used with coal-fired boilers since the handling properties of the wood pellets are comparable to coal.

Many power plants such as Drax and Fiddlers Ferry power stations in the United Kingdom as well as Avedøre plant in Denmark that co-firing of coal and wood pellets as an initiative to potentially reduce the carbon dioxide emissions (Henderson, 2015). In addition to that, the government of Netherlands also has the intention to mandate co-firing of biomass at all existing coal-fired power plants (Verhoest & Ryckmans, 2012). Besides co-firing, the use of biomass as the main fuels also been practised worldwide.
The increase in demand of the lignocellulosic biomass fuels in the energy sector can be witnessed in the growth of global wood pellet production between 2010 to 2014, which increased from 16 to 25 million tonnes as reported by Matthews, (2015). This growth could be triggered by the introduction of several European biomass standards such as EN 14961, which is for classification and specification of biomass in 2011 (Audigane et al., 2012). The lignocellulosic biomass has received increasing attention especially in the utilisation of the thermally treated products that enhanced its physical and thermal properties. The thermally treated products have properties that are better or equivalent to the coal. Gasification, combustion, pyrolysis and torrefaction are among widely used thermochemical applications that had been proven to upgrade lignocellulosic biomass quality. Besides that, pelletization also has improved the quality of the biomass fuels as well as eases the handling and transportation of the material.

However, similar to coal, biomass fuels are also prone to self-heating due to chemical oxidation when reacting with oxygen in the air during its storage period. The self-ignition can happen during storage if the heat generated by the chemical oxidation starts to accumulate and does not dissipate to the surrounding of the stored biomass fuels. The fire triggered by self-heating of stored bulk biomass fuels causes serious safety issues as it is difficult to detect when it started to occur. Besides the chemical oxidation, heating due to the microbiological activities can also happen in biomass fuels due to its properties such as hydrophilic nature and high moisture content.

The accidental fires caused by dust explosion and self-ignition can cause hazards to workers, storage structure as well as to the economic loss (Guo, 2013a). Serious injuries and sometimes fatalities are also witnessed in such fires. In addition to those, the undesired fires can cause emission of carbon dioxide and toxic gases such as carbon monoxide through incomplete combustion (Ferrero et al. 2009). Thus, prevention of the self-ignition is economical and safety-relevant issues that need to be studied and understood for accident prevention in the future.
There are many recorded accidental fires associated with the self-heating of biomass. For example, an estimated average of 14070 fires per year between 2005-2009 had been reported caused by self-ignition or chemical reaction in United State, with 20% of fires in storage properties were ignited by biomass material such as crops (Evarts, 2011). Data of major accidents associated with biomass energy production had been presented by Casson Moreno & Cozzani (2015), where 30% of the fires recorded was started by self-ignition of sawdust in stockpiles.

Table 1.3 provides examples of fire incidents associated with the self-heating of biomass during the last ten years in the United Kingdom. Regardless of many incidents of fires started due to the self-heating of biomass, there is still a lack of risk and safety awareness in this sector unlike in the coal industry. Thus, many researchers had suggested the need for further research in this area to fill in the gap. Moqbel et al. (2010) and Veznikova et al. (2014) have suggested studying the tendency to self-ignition, especially for wood-based fuels since their dangerous properties are insufficiently considered during handling and storage.

Table 1.3: Fire incidents associated with the self-heating of biomass in the United Kingdom for the past ten years

<table>
<thead>
<tr>
<th>Date</th>
<th>Place</th>
<th>Descriptions of events</th>
</tr>
</thead>
<tbody>
<tr>
<td>27th August 2016</td>
<td>Ancaster, Lincolnshire</td>
<td>Fire started at a conveyor belt in recycling plant</td>
</tr>
<tr>
<td>26th July 2013</td>
<td>Lawrence Recycling, Kidderminster</td>
<td>Self-ignition started from the heat generated in the recycling material</td>
</tr>
<tr>
<td>21st July 2013</td>
<td>Nature’s Choice, White Township</td>
<td>Blaze fire started from self-ignition of large pile of mulch</td>
</tr>
<tr>
<td>16th May 2013</td>
<td>Good 2 Grow, Beenham</td>
<td>Compost fire from self-ignition of wood mulch</td>
</tr>
<tr>
<td>12th May 2013</td>
<td>Todd Waste Management, Thirsk</td>
<td>Self-ignition from decomposition of organic material</td>
</tr>
<tr>
<td>18th September 2012</td>
<td>Arcwood Recycling Stanton-by-Dale</td>
<td>Blaze fire started from recycled wood</td>
</tr>
<tr>
<td>5th September 2012</td>
<td>Potts Farm, Farnham</td>
<td>Self-ignition caused by stored hay</td>
</tr>
<tr>
<td>27th February 2012</td>
<td>Tilbury Power Station, Essex</td>
<td>Fire caused by self-heating of wood pellets</td>
</tr>
<tr>
<td>5th November 2011</td>
<td>Port of Tyne</td>
<td>Fire started in a conveyor transfer tower storing biomass pellets</td>
</tr>
</tbody>
</table>
However, the main cause of the self-ignition of biomass fuel is so unique from one to another. It varies according to types of biomass and their physical, chemical and biological properties. Therefore, due to its wide possibilities of the cause, the fire is hard to predict and mitigate. However, the storage conditions of the biomass can be monitored and always have a high correlation with the heat generated by the system during the storage period. The heat generated due to self-heating depends on many parameters which can be categorised as a controllable, uncontrollable and conditionally controllable parameter (Arisoy & Akgun, 2000). Therefore, the focus of this study should always be on the approaches to prevent biomass fuels self-combustion during the storage period, where the storage conditions are considered as a controllable parameter. Thus, the critical storage conditions can be predicted to avoid the self-ignition.

Many studies have been done to predict the behaviour of the biomass material that has a tendency to self-heat in concealed storage facilities as a silo (Larsson et al., 2012; Malow & Krause, 2008; Guo 2013; Ramírez et al., 2010; Carrera-Rodríguez et al., 2011). However, there is limited research done on the tendency of self-heating of biomass fuels in open storage area. Over the past 20 years, many studies were done to investigate the self-heating behaviour, which measured the ignition kinetic parameters and the critical conditions that can lead to self-ignition (Chen et al., 2013). Hence, this study focuses on predicting self-heating process due to chemical oxidation by considering its kinetic parameters and the critical storage condition of the bulk biomass fuels in indoor open storage piles.

A recent positive development for replacement of coal by a sustainable source in the energy sector on a global scale had driven biomass fuel industry to produce biomass fuels that have similar properties to coal. Thus, many researchers had come out with methods to improve the energy density of biomass fuels, which will give added commercial value to the material produced. Among others, torrefaction is one of the proven methods to upgrade the quality of the biomass fuels. Besides that, the rationale behind the torrefaction process on wood pellets is to achieve a final product that is superior to the conventional wood pellets.
This study is focused on the self-heating behaviour of those two completely different materials, which are conventional wood pellets and thermally treated wood sample using microwave torrefaction technology. Ceballos et al. (2015) had pointed out several gaps in the study of torrefied biomass, which include the need to understand its self-heating propensity in stockpiles. Hence, this thesis is aimed to understand the self-heating behaviour of such biomass based on their physical and thermal properties that may influence the self-heating propensity in the stockpile. The study is carried out by simulating the self-heating behaviour of bulk biomass fuels based on experimentally determine kinetic properties. Several simulations were performed using COMSOL Multiphysics® software to obtain a deeper understanding of the effect of the controllable parameters on the self-heating behaviour of biomass fuels in piles during storage.

1.2 Approach of the research

The focus of this study is to examine the self-heating behaviour of the biomass fuels that undergo microwave torrefaction process. Microwave torrefied biomass has identifiable improvement in the energy density and the thermal kinetics parameters of the biomass; therefore it was used for this study. This study investigates the parameters affecting the self-heating behaviour of biomass storage. The findings from this study will particularly highlight the distinct changes of material properties before and after the microwave torrefaction process. It offers the opportunity to compare the reactivity of the biomass that undergoes the microwave torrefaction process and the non-torrefied biomass.

Furthermore, many studies on conventional torrefied biomass had reported various reaction conditions such as torrefaction temperature; inert gas and reaction time can lead to various solid, liquid and gaseous products. In addition to that, the higher torrefaction temperature or longer residence time resulted in torrefaction removes moisture, volatiles and degrades parts of its carbohydrate fraction, resulting in a product with higher energy density, better grindability and less moisture absorption (Nunes et al., 2013; Shang, 2012; Stelt et al., 2011).
Therefore, in this study the variation of process conditions for producing the torrefied sample was neglected and only torrefied sample produced using microwave process was considered in the self-heating propensity study. The study is focused on the variability of storage conditions that reflected the variable reactivity of both samples. There is a growing needs to determine whether the microwave torrefaction will increase the reactivity of the biomass and does the torrefied biomass needs to be handled in the same way as material that prone to self-heating.

This work also has no intention on examining which process condition that will contribute to a biomass fuel with higher reactivity by comparing the sample produces using conventional torrefaction method. Therefore, in this work, the comparison between the non-torrefied sample and microwave torrefied samples were adequate. The experimental work carried out to determine the chemical and physical properties, kinetic parameters as well as to examine the heating behaviour of both fuels. Information gathered from the experimental works is used in the simulation of the heating behaviour of the fuels in stockpiles.

The properties of the samples play a substantial role in the reactivity towards chemical reaction in air. Therefore, the main parameters associated with the self-heating phenomena were identified by the thermogravimetric analysis. The composition of moisture content, fixed carbon, volatile matters as well as ash is acquired from the proximate analysis using thermogravimetric analysis. While the composition of carbon, hydrogen, nitrogen and oxygen are obtained from the ultimate analysis.

Furthermore, the characteristic temperature observed from the thermogravimetric analysis can be used to measure the tendency of the material to undergo a self-heating process. Also, thermogravimetric analysis was done to obtain information of the reactivity of the samples as well as their kinetics in the air and the inert environment. Mass change measurements over temperature provide key information on the decomposition behaviour in the air and the inert environment.

Additionally, to examine the heating behaviour of the sample in bulk size, further analyses on the self-heating behaviour of the samples in a larger scale is in an oven-
controlled environment. The bulk oven test is significant to observe the temperature profile within the bulk sample in an oxidative environment at various oven temperature and bulk size. Besides that, the initial heating temperatures used for the non-torrefied sample were very similar to conventional torrefaction process which are between 180°C to 200°C, therefore, it can be considered that the torrefaction process was included in the oven test.

Table 1.4 shows all the experimental work done to determine the physical and chemical properties, kinetic parameters as well as the assessment of the thermal behaviour of samples.

Table 1.4: Experimental work done in the research

<table>
<thead>
<tr>
<th>Experimental tests</th>
<th>Purpose of the experiments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A. Sample properties determination</strong></td>
<td></td>
</tr>
<tr>
<td>i. Calorific value</td>
<td>Determination of the calorific value of the samples</td>
</tr>
<tr>
<td>ii. Proximate analysis</td>
<td>Determination of the moisture, fixed carbon volatile matter and ash</td>
</tr>
<tr>
<td>iii. Ultimate analysis</td>
<td>Determination of carbon, nitrogen, volatile matter and oxygen content</td>
</tr>
<tr>
<td><strong>B. Reactivity evaluation and kinetic parameters determination</strong></td>
<td></td>
</tr>
<tr>
<td>i. Thermogravimetric analysis</td>
<td>Reactivity evaluation based on characteristic temperature of the thermal decomposition</td>
</tr>
<tr>
<td></td>
<td>Determination of the kinetic parameters</td>
</tr>
<tr>
<td><strong>C. Large scale heating experiment</strong></td>
<td></td>
</tr>
<tr>
<td>i. Oven bulk test</td>
<td>Determination of heating behaviour of the samples heated up at various bulk sizes and oven temperatures.</td>
</tr>
</tbody>
</table>
1.3 Aim, Objectives and novel contribution

1.3.1 Aim and Objectives
This study is aimed at investigating the heating behaviour of microwave torrefied samples and non-torrefied sample to predict the self-heating of the fuels in stockpiles. The objectives of this research are:

(i) to determine the effect of physical and chemical properties of biomass fuels towards self-heating propensity using proximate and ultimate analysis;
(ii) to examine the thermal decomposition behaviour of fuels samples properties using thermogravimetric analysis;
(iii) to analyse the kinetic parameters of thermal decomposition of biomass fuels at a low heating rate;
(iv) to investigate the effect of bulk size and oven temperature on the heating behaviour of the biomass samples in oven bulk test;
(v) to develop a numerical model that can predict the effect of piles height and ambient temperature on self-heating behaviour of biomass samples; and
(vi) to validate the numerical model with the results from oven bulk test

1.3.2 Novelty of the research
The novel contribution of this research is the determination of the thermal behaviour of the microwave torrefied sample in comparison to the non-torrefied sample. Although a lot of researchers had been devoted to investigating the self-heating behaviour of biomass fuels, there is still very limited data on the self-heating behaviour of torrefied biomass fuels. In addition to that, to the best of the author’s knowledge, the study on self-heating behaviour of the sample prepared using microwave torrefaction has not been reported in any research, therefore requires further research. Previous studies on torrefied biomass fuels provided evidence that, the biomass fuel produced from microwave torrefaction increased the reactivity of the biomass. Therefore, additional precaution steps must be taken during storage and handling phase. The novel aspect of this research also established from the validation of the numerical model used in the simulations against the experimental data. In
brief, the findings from this research will help to examine the self-heating propensity of the biomass fuels produced using microwave torrefaction process.

1.4 Layout of the thesis

Chapter 2 presents the literature review on biomass usage as an alternative energy source and the fundamental theories of self-heating behaviour of the biomass fuels. Chapter 3 describes experimental method for characterisation of the samples used in this work, which include proximate and ultimate analysis. Chapter 4 covers the experimental work to evaluate the reactivity of the samples as well as their kinetic parameters. Also, the thermal decomposition behaviour of both microwaves torrefied and non-torrefied samples are presented. Chapter 5 presents the bulk tests to evaluate the reactivity of the samples in a larger scale. The relationships between the oven temperature and bulk size towards induction time are discussed. Chapter 6 examines the effects of geometry of the piles and the critical ambient temperature on the self-heating propensity using numerical simulation. Chapter 7 summarises all the experimental and simulation work in this thesis. Conclusions and recommendations for future work are presented.
Chapter 2 Literature review

This chapter presents a literature review of current research on fundamental behind the upgrading of lignocellulosic biomass using thermal treatment process for energy production and the self-heating behaviour of the biomass fuels in comparison to coal. This chapter also reviews the experimental work on self-heating evaluation.

2.1 Energy recovery from biomass

There are many alternative technologies available today that are able to recover the energy from biomass by converting biomass into solid fuel. However, the characteristic such as calorific value, moisture content as well as the amount of ash is unique for each biomass due to their differences in composition. Thus, it requires suitable conversion technology to convert the biomass into solid fuel efficiently.

There are two major technical approaches to convert lignocellulosic biomass into energy, which include thermochemical conversion and biochemical conversion. Thermochemical processes that widely used for biomass conversion process are pyrolysis, gasification and combustion, to produce heat or energy carrier such as charcoal and bio-oil. On the other hand, the biochemical conversion involved processes such as anaerobic digestion, alcoholic fermentation and aerobic biodegradation includes the process of a microorganism to break down the biomass into biomass-based energy fuels (Saxena, Adhikari, & Goyal, 2009; Balat, 2006). Figure 2.1 shows the possible route for thermochemical conversion of lignocellulosic biomass, while Figure 2.2 shows the possible route for the biochemical conversion.

According to Chiang et al. (2012), between these two types of technologies, thermochemical conversion has higher conversion efficiency with the better ability to break the organic compounds as well as shorter reaction time. Thus, thermochemical conversion is a favourable path for biomass concerning those advantages. The prior knowledge of the biomass such as chemical composition is necessary for choosing
the energy conversion process. The right biomass fuel is chosen based on its homogeneity, conversion technology and the plant design (Patel & Gami, 2012).

Figure 2.1: Possible routes of conversion for biomass fuels using thermochemical processes (Iowa Energy Center, 2016)

Figure 2.2: Possible routes of conversion for biomass fuels using biochemical processes (Badiei et al., 2014; Balat, 2006)
Pyrolysis is a thermochemical decomposition which occurs between 400°C and 900°C in the absence of oxygen. The pyrolysis breakdown of wood produces chemical substances that can be used as substitutes for conventional fuels. The biomass fuels can be pyrolyzed to produce liquid fuel or gases such as methane and hydrogen. On the other hand, combustion is an exothermic chemical reaction that produces substantial heat when the fuel reacts with air or oxygen (Tapasvi, 2015; Beever, 1995). The main steps in combustion process are drying, devolatilization, gasification, char combustion and the gas-phase oxidation (Garcia et al., 2012; Khan et al., 2009). According to Vargas et al. (2012), direct combustion is the most commonly used technology to convert biomass into heat.

Gasification is the conversion process of biomass to a gaseous fuel by heating in a medium such as air, oxygen or steam at high temperature (more than 700°C). The gasification process converts the intrinsic chemical energy of the carbon in the biomass into a combustible gas in two stages (McKendry, 2002). The gas produced from the process is more versatile to use compared to the raw biomass and can be utilised in the energy sector or used as a chemical feedstock to produce liquid fuels.

The energy source for the thermochemical process is usually solid biofuels such as a wood pellet or biochar. Therefore, to meet the energy demand, those solid biofuels need to be stored at the power plant prior to usage. For that reason, the need for safe handling and storage of such fuels is crucial to maintain continuous energy supply. Co-firing is widely used option for energy conversion, as it can utilise the existing coal power station without major modification. Compared to coal, biomass is bulkier and degrades easily as well as having properties that are unfavourable to coal. Therefore, with co-firing, the drawback can be improved.
2.1.1 Issues of biomass fuels as energy source

Several weaknesses of biomass need to be taken into consideration before using it as an energy source in the thermochemical process. Therefore, efficient use of the raw biomass can be achieved by transforming the biomass into fuels in solid, liquid and gaseous form via various pre-treatment processes.

Several concerns that had been reported associated with biomass are high moisture content, non-uniform physical properties, low calorific value, low energy density, structural heterogeneity, hydrophilic nature and low bulk density (Eseyin, Steele, & Jr., 2015; Nhuchhen, Basu, & Acharya, 2014; Sadaka et al., 2014; Järvinen & Agar, 2014; van der Stelt et al., 2011). In addition to that, the lignocellulosic biomass is reflected as thermally unstable by Prins et al. (2006a) even though having a low nitrogen, sulphur and ash content.

Biomass also inclines to be less dense than coal resulting in lower energy density. Low energy density is an unappealing aspect as it implies high transportation cost per energy unit as well as more storage area that leads to higher logistic cost (Nunes et al., 2014). Therefore, normally biomass fuel needs to undergo densification or palletization to increase its energy density. The ash content in biomass is also much higher than coal. The higher ash content reflected as a significant challenge for biomass combustion, due to the ash related problem during combustion such as slagging and fouling (Liu & Han, 2015; Deutmeyer et al., 2012; Darvell et al., 2010; Ciolkosz, 2010).

The latest review on the disadvantages of biomass compared to coal can be found in (Vassilev et al., 2015), where they pointed out that the despite the disadvantages of biomass in biofuel and biochemical application, the main environmental, economic and social benefits seem to counterbalance the shortcomings. Besides that, the drawbacks mentioned earlier can be resolved through the pre-treatment process of biomass (Chen & Kuo, 2011).
2.2 Pre-treatment methods for lignocellulosic biomass

The positive growth of sustainable energy sources such as biomass had driven biomass fuels industry to produce biomass fuels that have similar properties to coal. For example, as shown in Table 1.2, the energy density of wood is between 2.0 to 3.0 GJ/M$^3$ while the energy density of coal is between 18.4 to 23.8 GJ/M$^3$. These figures show that wood has a relatively lower energy density compared to coal. Therefore a pre-treatment is needed for wood to achieve a higher energy density that is comparable to coal.

Thus, a pre-treatment process is needed to upgrade the quality of the lignocellulosic biomass before being used as fuel in the energy sector. Pre-treatment is necessary for lignocellulosic biomass utilisation to modify the size, shape, and density of the biomass to match fuel specification or the thermochemical process. The upgraded lignocellulosic biomass had been proven suitable to be used for the co-firing with coal in the existing coal-fired power plant, without requiring major modifications to the plant (Liu et al., 2016). Besides that, it is essential to upgrade the lignocellulosic biomass fuel prior utilisation to ease its handling and transportation as well. In summary, the aims of lignocellulosic biomass pre-treatment had been recognised by Tapasvi (2015) to homogenise the biomass feedstock, increase the biomass energy density and improve biomass storage stability.

According to Demirbas (2004), the conversion of biomass via pre-treatment process is to transform a raw material, which is initially hard to handle, bulky and of low energy concentration, into a fuel with upgraded physicochemical characteristics that allows cost-effective storage and handling. Thus, pre-treatment for utilisation of the biomass for energy conversion is needed to address the shortcoming of the raw biomass. Harmsen et al. (2010) stated that the criteria lead to an improvement of lignocellulosic material after pre-treatment are (i) increasing the surface area and porosity, (ii) alteration of lignin structure, (iii) removal of lignin and/or hemicellulose, (iv) partial depolymerization of hemicellulose, and (v) reducing the crystallinity of cellulose. Table 2.1 summarised the purposes of each pre-treatment along with the examples.
However, the thermochemical pre-treatment pathway is more favourable to prepare the biomass for energy conversion due to the higher conversion energy as well as the conversion is faster compared to biochemical pre-treatment (Huang et al., 2016). Thus, the further explanation of thermochemical pre-treatment is presented afterwards as this study is focusing on the torrefied biomass fuels which are a product of thermochemical pre-treatment.

Table 2.1: Categories of pre-treatment of lignocellulosic biomass (Sindhu, Binod, & Pandey, 2016; Badiei et al., 2014; Harmsen et al., 2010)

<table>
<thead>
<tr>
<th>Categories</th>
<th>Purposes</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical pre-treatment</td>
<td>To increase the available specific surface area, and reduce the degree of polymerization and cellulose crystallinity</td>
<td>grinding, milling and pelletization</td>
</tr>
<tr>
<td>Chemical pre-treatment</td>
<td>Chemical reactions introduced for disruption of the biomass structure</td>
<td>liquid hot water, alkaline hydrolysis, acid hydrolysis, oxidative delignification</td>
</tr>
<tr>
<td>Thermochemical pre-treatment</td>
<td>To alter the chemical composition and physical structure of lignocellulose substrates</td>
<td>pyrolysis, gasification and torrefaction</td>
</tr>
<tr>
<td>Biochemical pre-treatment</td>
<td>To degrade hemicellulose and lignin for ethanol production using metabolite of a microorganism</td>
<td>degradation using white, brown and soft rot fungi to produce ethanol</td>
</tr>
</tbody>
</table>

2.2.1 Thermal pre-treatment

Various types of thermal pre-treatment available nowadays to treat lignocellulosic biomass. As mentioned earlier, to meet the demand of carbon neutral energy production, solid biofuel is preferable for co-firing with coal or entirely replace coal in power generation. Thus, to produce solid biofuels, the best option is to use thermal pre-treatment.
Thermal pre-treatment alters the chemical composition as well as the physical structure of the biomass. According to Bergman (2005), the combination of physical pre-treatment such as densification with the thermal pre-treatment like torrefaction is often proposed as an alternative to improve the physicochemical properties of biomass. Based on the various thermal pre-treatment, torrefaction is a promising technology to prepare the biomass for the further conversion process.

### 2.2.2 Torrefaction technology

Torrefaction has been acknowledged as one of the most assuring pre-treatments that can address some of the limitations discussed earlier by improving the quality of the biomass fuel. The process is intended to upgrade the lignocellulosic biomass into more homogenous fuel, which can be converted into energy. According to Prins (2005) and Gardbro (2014), the research on torrefaction process had started in France in 1930’s. However, there was no publication documented until 1984, and the first process patented in the late 19th century. Some researchers such as Deutmeyer et al. (2012), Melin (2011) and Lanigan (2010) had considered the torrefaction process to be similar to the technique used to roast coffee beans that had been practised since the late 13th century.

Torrefaction is similar to the pyrolysis, where both of the processes were carried out under the inert environment at atmospheric pressure. However, the main difference is the reaction temperature applied during the process. During the torrefaction, a lower reaction temperature applied to the process, which is between 200-300°C. Wet torrefaction is as treatment of biomass in a hydrothermal media or hot compressed water, at temperatures within 180–260°C (Bach et al., 2013). Dry torrefaction is the thermal treatment of biomass in an inert environment at atmospheric pressure and temperature within the range of 200–300 °C. The torrefaction discussed in this thesis is dry torrefaction technique unless stated otherwise.

The reaction temperature recommended by Prins et al. (2006b) is below 300°C, as they had discovered that a fast thermal cracking of cellulose happened when temperature above 300-320°C. The fast thermal cracking may cause tar formation.
Rousset et al. (2011) had divided the torrefaction process into two categories, which are light torrefaction (reaction temperature less than 240°C) and severe torrefaction (reaction temperature above 270°C). Thus, torrefaction sometimes referred as mild or partial pyrolysis.

Besides the reaction temperature, another distinct difference between torrefaction and pyrolysis is the maximisation of the solid yield, where the biomass weight reduced while the energy content is sustained. Thus, the maximisation of the solid yield where the biomass is converted to a carbon-rich solid product is the major motivation of torrefaction. The high solid yield can be achieved by removing the water and carbon dioxide from the lignocellulosic biomass, through the decomposition of biomass at the low heating rate, which is less than 50°C/min (Wang et al., 2012; Bergman & Kiel, 2005). As a result, the main objective of torrefaction is to remove oxygen from the raw lignocellulosic biomass. Therefore the product from torrefaction process is a solid biomass with a significant lower O/C ratio compared to its raw material (van der Stelt et al., 2011) and eventually increased its energy density.

### 2.2.3 Conventional torrefaction

The components in biomass play important roles in torrefaction process due to the different temperature effect on each component. The study of temperature ranges for peak thermal degradation of cellulose, hemicellulose and lignin, did by several researchers can be found in Nhuchhen et al. (2014). The temperature range presented had confirmed that hemicellulose degraded at a lower temperature range compared to cellulose, whereas lignin degrades at wider temperature range. Thus, the reactivity of the hemicellulose in biomass is the main attribute to the mechanism of torrefaction at the temperature range between 200°C-300°C. However, in that temperature range, cellulose and lignin are found to be less reactive.

According to Prins (2005), the kinetics of torrefaction reactions in temperature between 200°C to 300°C can be expressed by a two-step mechanism. While the first step is represented by the hemicellulose decomposition follows by the cellulose decomposition. The loss of bound moisture as well as thermal degradation to form
volatile products lead to reduced mass yield of torrefaction process (Prins et al. 2006a). Based on their research on torrefied willow, a large fraction of biomass solid is yield during the first step, which around 70-88% compared to only 41% for the second step. According to Ben & Ragauskas (2012), despite the biomass source, significant decomposition of hemicellulose happened at 240ºC, which explains the chosen reaction temperature for the torrefaction process that is below 300ºC, where the objective of the process achieved.

Subsequently, biomass material with a high hemicellulose content tends to have less solid product yield compare to the one with lower hemicellulose content (Nhuchhen et al. 2014). They also speculate that, within the torrefaction temperature range of 200-300ºC, the cellulose degradation might have enhanced by the water vapour and acids released from the hemicellulose degradation. As a result, the solid product from the torrefaction process has more lignin content, which is more stable component than hemicellulose and cellulose.

Torrefaction process leads to devolatilization of the cell wall constituents, which will cause enlargement of the lumina and create more available space for free water to occupy (Kymäläinen et al., 2015). Hence, torrefaction had been known for its advantages to improve the energy density and shelf life of the biomass. However, due to the incomplete devolatilization during torrefaction, torrefied biomass produced contained some volatile matters. In study done by Tumuluru et al. (2011) they had listed eight (8) parameters that influence the torrefaction process, namely (a) reaction temperature, (b) heating rate, (c) absence of oxygen, (d) residence time, (e) ambient pressure, (f) flexible feedstock, (g) feedstock moisture and (h) feedstock particles size. According to van der Stelt et al. (2011) torrefaction process can be divided into stages of the temperature-time as presented in Figure 2.3:
The reaction occurred during the torrefaction process demonstrates a positive outcome to the chemical and physical properties of the raw biomass, which changes its properties closer to those of bituminous coal (Nunes et al., 2014). They had established that biomass torrefaction can produce a torrefied biomass with an energy density that is almost equal to coal but denser than in wood. The properties of the solid biomass products fall between the properties of coal and biomass (Fisher et al., 2012). On the other hand, Rousset et al. (2011) proved that the characteristics of the torrefied bamboo tend to fall toward those of low-rank coals. The same findings can also be found in Tapasvi (2015), where the composition of the torrefied Norwegian birch and spruce were closer to coal with lower volatile matters and higher carbon content. Thus, the torrefied solid products can be used to substitute charcoal in some applications such as fuel for domestic cooking stoves, residential heating, and manufacture of improved solid fuel products such as fuel pellets or barbeque briquettes for commercial and domestic uses.

There are many types of reactors available nowadays, and each reactor has its distinct properties and is unique to certain biomass physical and chemical
characteristics only (Medic, 2012). The overview of torrefaction reactors with different technology by several developers in Europe, North America, and Brazil can be found in Shang (2012), which consists of several categories such as conveyor reactor, moving bed reactor, rotary drum reactor, fluidised bed reactor and microwave reactor. All the categories mentioned earlier use electrical heating method except for microwave reactor that uses microwave radiation to heat up the sample. The method of torrefaction using electrical heating is referred as conventional torrefaction in this thesis. Subsequently, this study is focused on the torrefied biomass fuels using microwave reactor.

2.2.4 Microwave torrefaction

The general definition of microwaves is electromagnetic waves, which frequencies fall between 300 MHz and 300 GHz with corresponding wavelengths between 1m and 1mm, respectively. Figure 2.5 shows two mechanisms of microwave heating that contribute to the heating of the materials quickly and uniformly are; dipole rotation (dipolar polarisation) and ionic conduction (Anwar et al., 2015; Lanigan, 2010). The dipolar rotation happened when the polar molecules such as water in microwave field rotate to align themselves with the changing field. The friction and collision between a dipole molecule and the surrounding molecules dissipate heat. Whereas, the ionic conduction occurs when the free ions oscillate back and forth under the electric field caused by the microwaves energy. The ionic motion causes the rapid heating of the material.

Figure 2.4: Mechanism of microwave heating (Anwar et al., 2015; Lanigan, 2010).
Thus, unlike conventional heating that transfers heat to the materials, microwaves produced heat by converting the electromagnetic energy to the thermal energy (Wang et al., 2012). However, due to those heating mechanisms, only materials with the dielectric properties can absorb microwaves. According to Huang et al. (2016); Gronnow et al. (2013); Wang et al. (2012); Lanigan (2010), the microwave chemistry suggests many advantages of the microwave torrefaction over the conventional torrefaction. Some of the benefits discussed are: the shorter reaction time compare to the conventional method and does not need direct contact between the heated material and energy source, selective heating and energy saving.

One of the main advantages discussed by many researchers is the mode of energy transfer when heating using microwave radiation over the conventional heating. The microwave energy is delivered directly into the material through molecular interaction with the electromagnetic field. While for the conventional heating method, the heat is transferred into the material via convection, conduction and radiation from the surfaces of the material. Therefore, Lanigan (2010) explained that the heating using the microwave is an energy efficient process as the microwave irradiation provide volumetric heating throughout the material while the conventional heating heats the material that in contact with the reaction vessel before the whole volume. However, main parameters that affect the torrefaction reaction are microwave power and reaction time.

### 2.2.5 Torrefied biomass fuels

The solid product from the torrefaction sometimes referred as biochar or simply torrefied biomass. Nunes et al. (2014) identified the improvement of the physical properties biomass torrefaction such as the grindability, the particle shape, size, and distribution as well as improves the pelletability. The torrefied biomass has lower volatile matters and oxygen, a higher calorific value as well as higher ash and carbon content (Huang et al., 2012; Jones et al., 2012; Wang et al., 2012; Arias et al., 2008). The characteristics mentioned earlier were more similar to that of coal, which makes the torrefied biomass suitable for co-firing with coal in power plant.
Given the high volumetric energy density, the solid torrefied biomass has been recognised to be suitable for combustion or gasification application in commercial and residential (Tumuluru et al., 2011) or as co-firing material in power plant (Nhuchhen et al., 2014). Besides the chemical properties, torrefaction process also upgrades the physical properties of the biomass. For example, the study by Li et al. (2012) reported a decreased of water adsorption capacity after torrefaction of wood material. In addition to that, the study also mentioned that moisture content reduced to almost half of its raw material.

Despite all the benefits presented earlier, due to the brittleness of the torrefied biomass, Michael Wild (2016) and Bergman (2005) claimed that more dust would be produced during handling compared to coal and extra care shall be taken to avoid dust accumulation as it said to be more reactive than coal dust. In addition, off-gassing issues in torrefied biomass showed an alarming risk that required mitigation by proper design of storage facilities along with safe handling procedures (Tumuluru et al., 2015).

The torrefaction process also changed the reactivity of the raw biomass. Researcher such as Liu et al. (2016) and Fisher et al. (2012) reported the decreases of biomass reactivity after torrefaction. However, the reactivity is affected by the torrefaction degree as well as the heating rate. The torrefied biomass is more reactive when the torrefaction process uses higher heating rate than the one produced in lower heating rate. The reactivity might have a good implication on the utilisation for combustion, but there is a possibility of the self-heating problem.

2.3 Storage of biomass fuels

The growing interest in the production of solid biofuels has led to the need for the storage in large amounts prior usage. The production of solid biofuels takes place all over the year. However, the higher demand for the solid biofuels mostly during the winter season due to the needs of energy for heating. Therefore, there is a necessity to store the solid biofuels for the long term. Nevertheless, there is also a need for short-term storage of biomass at the site.
To be able to operate continuously, a power plant needs to be able to handle large quantities of fuel. Therefore, delivery just in time prior usage is impossible. Consequently, the plant needs to have a buffer storage of fuel at the site to cope with the fluctuation demand, as well as delays in the delivery chain (Eriksson, 2011). The solid biofuels storage needed to be first in first out basis to avoid the degradation of the fuels during storage.

Current practice to store conventional wood pellets is usually in the silo to protect them from exposure to moisture, which can cause swell and disintegrate. Based on Biochar.net (2012), current practice for transportation and storage purposes of biochar is packed in a sealed 200 L drums or one cubic meter sacks. However, there is a possibility for the biochar to develop heat once exposed to air.

While for torrefied biomass fuel, the outdoor pile storage is not recommended as the best practice in stockpiling (Järvinen & Agar, 2014). Therefore, the study had suggested that storage practice for torrefied biomass fuel been treated similarly to wood pellet. The suggestion had been proven as a suitable storage practice by Kymäläinen et al. (2015), through their study of five-month storage trial of several types of biomass fuel including torrefied biomass. The findings from the study suggested that even though torrefied biomass fuel is hydrophobic, the liquid water can still be absorbed into the bulk through capillary action. However, the study recommended that the usage of the silo to store the torrefied biomass fuel be unnecessary, unlike the untreated pellet. Thus, roof-covered storage areas or simple covers are suggested as a cheaper option for storage condition of torrefied fuel, and this suggestion matches the recommendation in Koppejan et al. (2012). There are limited amounts of experimental data on the self-heating of torrefied biomass stored in large piles or heaps, but it most likely will follow the same self-heating mechanism of coal piles.

2.4 Self-heating leading to ignition

Self-ignition is a thermal runaway resulting from the self-heating process. Self-heating is the process where the materials achieve temperature higher than surrounding temperature due to the internal exothermic reactions. According to
Bowes (1971), the thermal explosion analysis by Semenov (1940) and Frank-Kamenetskii (1955) are the basis of the theoretical study of the self-heating and self-ignition in a stockpile. The phenomena of the self-ignition in a stockpile are described by two main parameters, which is the heat gained by the reaction and the heat lost to the surrounding. Self-ignition is a phenomenon where thermal runaway occurs due to self-heating of the material resulted in a transition from slow internal exothermic reaction to a rapid oxidation without external heat source such as spark or flame (van Blijderveen, Bramer, & Brem, 2013). Self-heating, on the other hand, is part of the self-ignition process that happened before the ignition occurs (Fan and Dong, 2011).

The imbalance between the heat produced by the reaction and the heat release through the stockpile may lead to the self-heating. Self-heating happens when the temperature of the stockpile is higher than the ambient temperature due to the internal exothermic reactions. The self-ignition in stockpiles is a thermal runaway resulting from the self-heating process that cannot be balanced by heat loss in the system. Self-ignition also commonly being termed as spontaneous ignition, auto-ignition and auto-combustion.

The mathematical model of self-ignition from the theory is based on the partial differential equation of heat conduction, where the governing equation of heat transfer is as shown in Eq. 2.1.

\[
\frac{\partial T}{\partial t} = a \left[ \frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} + \frac{\partial^2 T}{\partial z^2} \right] + \frac{Q(T)}{\rho c} \quad (\text{Eq. 2.1})
\]

Q is the heat source, W/m³, that is represented by an Arrhenius model as presented in Eq. 2.2

\[
Q = q \rho Ae^{\frac{-E}{RT}} \quad (\text{Eq. 2.2})
\]

Also, Figure 2.5, describes the behaviour of the system based on the heat balance due to the heat of reaction and heat loss. Based on the diagram, the thick curve line is the heat of reaction \(Q_{\text{react}}\) that increases in a non-linear form with the temperature. While heat loss to the surrounding is represented by, the three grey parallel lines, which are at the initial temperature, Ta. The three parallel straight lines marked a, b
and $c$ are the convective heat transport from the fuel bed to air stream for three primary air stream temperatures $Q_{\text{conv}}$. These lines cross the x-axis at air temperature ($T_o$). When $Q_{\text{conv}}$ is negative, the fuel bed is heated by the primary airflow.

![Diagram](image)

**Figure 2.5:** Heat produced by the reaction ($Q_{\text{react}}$) and the heat lost or gained due to convective heat transfer for three values of $T_o$ (van Blijderveen et al., 2010)

There are two essential parameters to assess the self-heating behaviour that lead to ignition; namely spontaneous ignition temperature and induction time. The spontaneous ignition temperature is defined as the highest surrounding temperature where no ignition occurs at a given volume, however the thermal runaway can happen if the surrounding temperature is higher than the self ignition temperature (SIT) where the heat released by the reactions inside the material is larger than the heat transmitted to the environment (Ferrero et al., 2008).

In the other hand, induction time or ignition delay time is the time taken by the material to ignite, which according to Saddawi et al. (2013) is a function of surrounding temperature of the bed where the self-heating and ignition take place. The difference arises in the induction time of material had been justified by Thomas & Bowes (1961) as a result of time taken of the material to form the active product material or to destroy the reaction inhibitor of the material.

The relationship between SIT and ignition delay times can be examined by the method in BS EN 15188:2007 (BS 15188, 2007). As tested by Saddawi et al. (2013), the results obtained from the test are a function of the size of the sample, where the
SIT will be lower for larger sample size and a longer time is taken to ignite (longer induction time).

### 2.4.1 Self-heating behaviour of coal

However, coal stockpiles are susceptible to self-ignition particularly if the large amounts are stored for long periods. The subtle change in the physiochemical properties upon weathering is called oxidation. Based on Nalbandian (2010) the accumulation of heat due to the heat generated from the oxidation process can lead to self-ignition. The oxidation process is affected by three ways of how oxygen affects the process (Veznikova et al., 2014). The three ways are physical adsorption of oxygen the, reaction of oxygen on highly active coal centres, and the chemical reaction of oxygen on coal. Thus, the mechanism of self-ignition results from the physical and chemical reaction of coal.

The process of self-ignition coal can be divided into four phases as listed below:

- **a.** Chemisorption of oxygen along with the increase of weight; from ambient temperature up to 70°C
- **b.** Initial release of oxidation reactions products and inner water; temperature range between 70°C to 150°C
- **c.** Production of larger amounts of oxidation reactions products; temperature range between 150°C to 230°C,
- **d.** Fast burning including production of soot; temperature above 230°C

Some factors that influence the process of self-ignition are chemical reactivity of the coal towards oxygen, atmospheric conditions, the moisture content of the coal, a method of storage, particles size, and physical properties of coal (Nalbandian, 2010; Schmal et al., 1985). The works by Beamish et al. (2000) show that the self-heating rate of freshly exposed coal to the atmosphere is higher than the aged coal. The rate of the rate is lower due to the loss of the reactive site for oxidation. Besides that, the propensity of self-heating is different between types of coal. Low-rank coals exhibit strong self-heating tendency compared to the matured coal (Ceballos et al., 2015; Fei et al., 2009; Hull et al., 1997; Krishnaswamy, Agarwal, & Gunn, 1996; Jones & Vais, 1991). Coal self-heating propensity is associated with coals containing high
volatiles (low-rank) compared to low volatile and high fixed carbon coals (high-rank). The volatile matters in coal decreases as the ranking increases.

### 2.4.2 Self-heating behaviour in biomass fuels stockpile

The temperature rises of biomass fuel in a stockpile occurs primarily due to the self-heating generated by chemical, biological or physical processes. Chemical self-heating takes account of the reaction between the gas and solid material such as oxidation. While for the biological self-heating, the metabolism of microbes within the biomass generates heat. Lastly, the self-heating concerning physical process includes effect related to moisture transport (Leslie, 2014). When storing large amount of biomass fuels, there is a high risk of self-heating occurrence in the stockpile due to a combination of those self-heating processes.

Biomass fuels have a potential to absorb oxygen during storage period to produce exothermic reactions in the same manners of risk is observed in coal storage period (Mateos et al., 2013). The oxidation at low temperature (below 200°C) can be observed for biomass fuels, where the oxidation rate depends on the surface area of the active site. Morrison & Hart (2012) had acknowledged the risk of self-heating processes in piles of biomass, which may be under the oxygen-limited environment and access to oxygen able to increase the heat generation and onset of a smouldering or thermal runaway.

The exothermic reaction can take place under various conditions depending on the materials observed. According to (Ramírez, García & Tascón, 2010), the initial stage of temperature rise in biomass stockpile happens typically due to the biological activity of microorganisms such as bacteria and moulds. The temperature of the system commonly rise to 55°C, and in some cases, the temperature may up to 75°C. Jirjis (2005) found the relationship between the fungal activities and the heat development in fresh willow shoot storage, where the spore counts increased was closely related to the heat development. Respiration is one of the processes that happened in all organic materials that were stored in silos or storage facilities as being explained by Ramírez et al. (2010). Respiration process will break down carbohydrates, proteins and fats into carbon dioxide, water and energy. The self-
heating process will happen when the energy is used by the cell to fuel the metabolic process which will release heat; if there is insufficient heat exchange between the reaction system and the external environment the temperature will eventually rise therefore will further increase the rate of reaction which then the product becomes even warmer.

Afterwards, the heat generation due to chemical oxidation takes place, and the heating up can occur up to at least 150°C. The same temperature pattern as explained before can be witnessed in the study by Hogland & Marques (2003) on the storage of waste pile. The study explained that the physical process due to adsorption or condensation might lead to heat generation. However, it happened at a low temperature (below 20°C), which is not significant in this work. Also, Thomas & Bowes (1961) highlighted the complexity of the self-heating process which may include more than one exothermic reaction, however in the first stage of reactions, only low heat generated is witnessed, which will not result in the ignition.

According to Krause (2009), there are four main criteria must take place for the spontaneous combustion to occur. The main criteria are: (1) the material must exhibit self-heating properties, (ii) the reaction that contributes to self-heating in the system must rapidly accelerate to reach high temperature until it extends to a thermal runaway, (iii) the thermal runaway phase must start a sustained smouldering, (4) the
sustained smouldering must spread to the outside surface of the material and possibly erupt into flaming. Figure 2.3 shows the phases concerning the criteria lead to the occurrence of a self-ignition process.

Meanwhile, Nelson et al. (2003) described the process of heat generation involving bulk organic materials into two stages of temperature only: (i) at low-temperature heat generated is due to the process of growth and respiration of microorganisms and (ii) at high-temperature heat generated is due to chemical oxidation of cellulosic materials. Their work also dismissed the heat generated due to a physical process. Therefore, this study is focused on the source of heat generation at high temperature; that only consider the chemical reaction of the biomass fuels without the effect of microorganism activities. Besides that, the samples used in this study are considered having a very low moisture content, which is an unfavourable environment for biological activity by microorganisms.

2.4.3 Factors influence the self-heating propensity

There are several factors, which have influence the tendency of self-heating of biomass fuels during the storage period. Two types of factors influence the self-heating process in biomass fuels during the storage period are controllable and uncontrollable factors. Controllable factors take account of the management of the pile condition itself, for example, the pile height. In the other hand, the uncontrollable factors include the characteristic of the biomass fuel. However, this study only focuses on the controllable factors that can influence the propensity of the biomass fuels self-heating.

Nalbandian (2010) had discussed some factors that influence the spontaneous combustion of coal, which included air-ventilation of the stockpile, silo or bunker, the atmospheric conditions of the storage area, the coal quality, coal moisture content and particle size. Hogland & Marques (2003) listed several factors that influence the temperature development in a waste storage pile such as particle size, the amount of organic material, moisture content, the size of the waste pile and the surface area of the waste fuel available to the reaction. Similarly, these factors also play roles influencing the self-ignition during storage of biomass fuels. Storage time,
surrounding conditions, species composition, form of the biomass, as well as geometry and structure of the storage pile influenced tendency of the biomass characteristics changes during storage period (Krigstin & Wetzel, 2016), which eventually affect the propensity of the self-heating to occur.

The compaction of the bulk material can also increase the propensity of the material to self-heat, due to the increase in the thermal conductivity as reported by Mateos et al. (2013) on the study of several types of lignocellulosic biomass with a particle size of 0.5 μm and 0.9 μm. While more uniform particle size of wood chips lead to higher compaction, which will reduce the heat dissipation to the surrounding especially in higher chip piles (Jirjis, 2005).

Pauner & Bygbjerg (2007); Wilen Carl & Rautalin (1993) concluded in their research that the ambient temperature of the storage area and the characteristic of the biomass fuel play an important role influencing the propensity of the self-heating. In addition to that, Guo (2013) considered the influence of the ventilation in the storage of wood pellets in a silo, which indeed a very significant factor to be taken into attention.

Many studies agreed that the stockpile geometry is a critical factor to be taken into account when storing biomass fuels in a bulk quantity (Murasawa et al., 2012; Ferrero et al., 2009; Li, Koseki, & Momota, 2006; Jirjis, 2005). The mechanism of the heat dissipated for from the stockpiles is through the surface even though the heat is produced in the whole bulk volume. Therefore, if the heat loss to the surrounding through the surface is unable to balance the heat generation, it can initiate the self-ignition.

For instance, Li et al. (2006) had established the relationship between the heat generation and the volume of the bulk pile. The heat generation in a pile is proportional to the its volume, while the heat loss is proportional to the surface area. Hence, volume to surface area ratio (V/A) has a strong influence on the self-ignition occurrence as the rate reaction associated with it. According to Cruz Ceballos et al. (2015), the higher surface area can be translated to more active sites. Thus, to be able
to control the influence of the geometry toward self-heating propensity, the height of piles is a controllable factor that can be managed during the storage period.

2.4.4 Biomass fuels ignition preventions

Many researchers had study the safe storage of biomass fuels in stockpiles regards physical and chemical properties that likely to cause self-heating. Table 2.2, listed the several countermeasure suggestions of the research based on the material studied. Based on the several examples of countermeasures suggested, the storage environment plays an important role to avoid the self-ignition occurrences. The surrounding temperature of the storage is a key factor to be taken into account along with the pile condition such as its height. Besides that, the physical condition of the stockpile such as the need to compact or bale the material can avoid the self-ignition. Thus, those factors are the focus of this study, where the effect of pile height, temperature and material porosity is presented.

2.5 Experimental study of self-heating behaviour

There are many analysis methods available to evaluate the self-heating propensity of the biomass material. Among the experimental techniques adopted by researchers to analyse the self-heating in biomass fuels are based on the one used for coal. The study of the self-heating of coal in the mining industry is well understood compared to the biomass fuel industry (García-Torrent et al., 2012). Similar to coal, the understanding of the kinetics of the biomass fuel oxidation is beneficial for the understanding, design and modelling of the industrial processes (Sima-Ella, Yuan, & Mays, 2005). The self-heating analysis can be categorised based on the method used to obtain the self-heating parameters, which characterised the material’s reactivity. The main kinetic parameters that influence the reactivity of the biomass fuels are activation energy and pre-exponential factor. Sima-Ella et al. (2005) elaborated that the activation energy affects the temperature sensitivity of reaction rate, while the pre-exponentials factor associated with the material structure.
### Table 2.2: Prevention suggestions from past studies on self-heating of biomass material in stockpiles

<table>
<thead>
<tr>
<th>Sources and biomasses studied</th>
<th>Suggestions</th>
</tr>
</thead>
</table>
| **Graham (2015)**            | - Long term storage of thermally treated wood pellet in an open barn with cover  
                             | - Short term option of outdoor storage for thermally treated wood pellet  
                             | - Storage of untreated white wood pellet in a fully enclosed environment  
                             | - Sloped edge pile |
| - Freshly harvested willow   |             |
| - Thermally treated wood pellet |               |
| - Untreated white wood pellet |               |
| **Mateos et al. (2013)**    | - Compaction of the stored material reduced the self-heating propensity; due to the reduction of air accessibility. |
| - Several types of milled lignocellulosic biomass (size: 0.5μm & 0.9μm) |             |
| **Guo (2013)**              | - Smaller silo size for storage system  
                             | - Increase the pellet age before storage  
                             | - Ventilation system inside the silo |
| - Wood pellet |             |
| **Koseki (2012)**           | - The removal of heat using a tube that is penetrated into the pile  
                             | - Active monitoring of the temperature and gas emission from the pile  
                             | - Reduce fermentation by covering the low part of the pile’s slope with incombustible material. |
| - Wood chips  
| - Chicken dung  
| - Refuse paper and plastic fuel  
| - Organic rubble  
| - Soy sauce squeezing residue |             |
| **Murasawa et al. (2012)**  | - Limitation of the pile height  
                             | - Adjustment of the surrounding temperature |
| - Soy sauce squeezing residue  
| - Fish meal residue |             |
| **Fu et al., (2006)**       | - Monitoring of the relative humidity and temperature inside bulk piles  
                             | - Avoid the long-term storage |
| - Refuse derived fuel  
| - Meat bone meal |             |
| **Hogland & Marques (2003)** | - Baling of the material to reduce the porosity  
                             | - Ensure pile is lower than 5m  
                             | - Construction of the wind barrier to protect the pile from strong wind |
| - Refused derived fuel  
| - Unsorted industrial waste |             |
2.5.1 Thermogravimetric analysis of thermal decomposition

Critical conditions for self-heating in stored biomass materials can be evaluated by using suitable laboratory-scale test method including thermogravimetric test, differential scanning calorimeter, differential thermal analysis, adiabatic calorimeter and isothermal oven tests using basket heating methods (Cruz Ceballos, Hawboldt, & Hellleur, 2015; García-Torrent et al., 2012). The thermal analysis such as thermogravimetric and differential scanning calorimeter require smaller sample size as well as less time-consuming.

Among the techniques used, the most common approaches are using thermo-analytical analysis, which based on the net amount of heat released by the sample as well as the dependence of certain parameters to the temperature (Avila, 2012). This technique including thermogravimetric analysis (TGA), differential thermal analysis (DTA) and differential scanning calorimeter (DSC) are widely used in self-ignition researches.

A thermogravimetric analysis is extensively used to determine the kinetic parameters for the sample as well as to evaluate its thermal stability. TGA was performed to measure the mass changes of the sample related to its temperature changes which provides the characteristic temperature of the sample. Many researchers such as (Jones et al., 2015; Della Zassa et al., 2013; Avila et al., 2011; Saddawi et al., 2010) used this method in the study of reactivity biomass materials. Nevertheless, this technique had been widely used in the study of coal as well.

Therefore, the significant parameters examined based on thermogravimetric analysis in term of self-heating behaviour as reported by Ramírez et al. (2010) are are the temperature maximum weight loss (TMWL), temperature of initial combustion (TIC) and characteristic oxidation temperature ($T_{charac}$).
The definition of each parameters are as below:

**a) Temperature of maximum weight loss (TMWL)**
TMWL is used as an indication of the reactivity of the sample tested. The maximum weight loss is due to the yield of volatile matter produced during the thermal degradation process. This value implies as an index of the reactivity of the samples. The higher the temperature reaches, the lower the reactivity of the sample.

**b) Temperature of initial combustion (TIC)**
TIC shows the temperature at the beginning of the combustion profile, which indicates that the combustion starts. The lower the TIC value, the lower the reactivity of the sample. This value will give a clear view on which temperature the material begins to combust and shows that if the material reactivity is higher in the air, it will start to combust at a lower temperature.

**c) Characteristic oxidation temperature (T_{charac})**
T_{charac} is a single oxidation temperature obtained when TG analysis was done in an oxygen environment. T_{charac} can be assign when using oxygen because the oxidation reaction takes place quickly and the point where the sudden weight loss can be determined. The relationship between T_{charac} and activation energy can be used to evaluate the classification of the samples concerning their ignition risk (Ramírez et al., 2010).

Differential scanning calorimeter (DSC) measures the temperature difference between the sample and a reference, which recorded against the temperature of the oven. Therefore, the exchanges of heat in the sample can be determined. DSC provides very sensible measurements with a small amount of sample (mg). According to Li et al. (2014) thermo-analytical technique using DSC is widely used to measure heat effect during coal oxidation.

The reactivity of the material in term of self-ignition study can be described as the propensity of the material towards rapid self-heating. Thus, the reactivity of the material can be examined based on how the material responds when being heated during TG analysis. Fisher et al. (2012) had emphasised that, even though there is no
accepted definition of char reactivity, the global char reactivity can be determined from mass loss histories, by defining it as the time derivative of the conversion. Therefore, this study adopted the same definition of char reactivity as earlier researchers.

However, thermal analysis operates on the relatively small sample (mg samples), which raise the issues in the sample representative towards actual heating behaviour as well as it reproducibility (Della Zassa et al., 2013). Therefore, the need of larger scale investigation is essential. Thus, the isothermal oven test is one of the preferable option.

2.5.2 Isothermal oven test

The isothermal oven test is conducted to investigate the larger scale of heating behaviour on the combustible material. The effect of temperature well as the volume over the self-heating propensity can be quantified using this technique. Besides that, the kinetic parameters of the heating process can also be found.

Three different curves can be obtained from this experiment. The curves are presented in Figure 2.7. The curve A is a subcritical curve where at chosen experimental temperature $T_A$ the sample becomes hotter towards the oven temperature. However, no exothermic reaction within the sample, thus no ignition is observed. While for curve labelled B, is the critical curve. At this temperature profile, the temperature of the sample slightly exceeds the oven temperature, $T_B$. However, it just surpasses for a slight time then tends to reduce towards the oven temperature. Lastly, if curve C is observed during the experiment, ignition will be detected. The supercritical curve observed when the heat generation in the sample surpasses its heat losses. Eventually, non-stationary conditions reached, and the sample temperature escalates rapidly over that of the oven ($T_C$), and ignition occurs.
The isothermal oven test uses crossing point (CP) method to find the kinetic parameters. This method assumes that when two temperature points inside the container are identical, there will be no conductive heat transfer between two points, leads to the heat transfer based on a transient solution as Eq 2.3.

\[ \rho C_p \frac{\partial T}{\partial t}_p = \rho \cdot \Delta_r \cdot h \cdot A e^{-\frac{E_a}{RT_p}} \]  

(Eq. 2.3)

Where the subscript \( p \) represents the condition at the centre of the sample. The identical temperature between this two location points is defined as the crossing point, where activation energy \( (E_a) \) can be calculated after \( T_p \), which is the ‘crossing point temperature’ and \( \frac{\partial T}{\partial t}_p \), is the temperature increase rate being examined experimentally. The activation energy can be calculated from the slope of \( \ln \left( \frac{dT}{dt} \right)_p \) against \( \left( \frac{1}{T_p} \right) \).
2.5.3 Numerical modelling of self-ignition

Various level of complexity to predict the condition of which biomass can undergo self-heating, starting with one-dimensional (1-D) to the more complex like two-dimensional (2-D) and three-dimensional (3-D). However, Fierro et al. (2001) had highlighted the disadvantage of time-consuming of the computational time with the possibility for the inaccuracy of prediction.

Many mathematical modelling had been done to study the self-heating process leading to ignition. Researchers manage to find the relationships between critical parameters to predict the possibility of ignition occurrence as been done by Ferrero et al. (2008); Ferrero et al. (2009); Lohrer et al. (2005); Fan & Dong (2011) and Pauner & Bygbjerg (2007).

The one-dimensional model equations are preferable, as it is less time consuming compared to two-dimensional (Arisoy & Akgun, 2000; Hull et al., 1997). However, the one-dimensional model will give less accurate results. Thus, many recent studies had run the simulations in two-dimensional as well as included complex coupled reaction such as biological and chemical oxidation in the numerical model in order to get more accurate and realistic results (Escudey et al., 2011; Luangwilai et al., 2010; Ferrero et al., 2008; Krause, Schmidt, & Lohrer, 2006; Akgun & Essenhigh, 2001). Also, Akgun & Essenhigh (2001) recognised the need of using the one-dimensional model as a first step approximation, but the model is incomplete and unrealistic. Therefore, a two-dimensional model is the simplest model that can give the accurate and realistic result. In the other hand, for the case of the prediction of critical conditions, the larger geometry managed to predict the critical conditions using the global developed model.

In order to for the parameters to be used in the simulation, thermal properties of the material can be examined using the thermal analysis such as thermogravimetric (TG) and differential scanning calorimeter (DSC). The temperature from the test will be applied in the numerical equation to determine the self-heating kinetic. Besides that, the prediction using numerical simulation can also be validated by experimental methods. Atreya (1998) had mentioned that there are three requirements in
experimental investigation of the ignition phenomenon, which are (i) a heat source to simulate the fire heat flux, (ii) a technique of mounting and exposing the sample to the heat source, and (iii) in case of piloted study, the initial source to ignite the sample. Numerical simulations will help in closing the gap when there’s a restriction in size or geometry of materials that need to be tested and often be used to complement extensive experiments (Lohrer et al., 2005).

2.6 Conclusions of the literature review

- The growth of demand for biomass fuel increased due to the awareness of using the carbon neutral fuel source for energy production. Therefore many new technologies to produced biomass fuels had been developed.
- The torrefaction technology managed to upgrade the quality of the biomass into a solid biofuel with high energy density and suitable for combustion or gasification for energy production. The torrefied biomass also a carbon neutral fuel, which is preferable compared to coal.
- The properties of torrefied biomass are comparable to the one in low ranking coal, with lower volatile matters and oxygen and higher calorific value, ash and carbon content.
- The torrefaction process changes the reactivity of the biomass based on the degree of torrefaction. Therefore, the torrefied biomass also likely to have issues of self-heating due to higher reactivity.
- Microwave torrefaction has advantages over conventional torrefaction with overall efficient process as well as promising technology application.
- Self-heating behaviour can be observed once the heat generated by the exothermic reaction is more than the heat losses to the surrounding. The self-heating leading to the ignition is a problem during the biomass bulk storage.
- The parameters affecting self-heating propensity of biomass fuel in bulk storage include ambient storage temperature, species composition, geometry of the storage (pile height), storage ventilation, physical and chemical properties of the biomass, surface area as well as the duration of the storage.
- Thermogravimetric analysis is a useful method to find the thermal kinetic for the biomass decomposition as well as the proximate analysis.
Chapter 2: Literature review

- Larger scale experiment is suitable to give more in-depth observation and measurement of the sample reactivity based on the effect of temperature and volume of samples towards heating propensity.
- Numerical modelling is a decisional-support tool that can help to simulate the heating behaviour on a large scale as well as to complement the findings from the experiment.
- Two-dimensional geometry model proven to have better advantages over one-dimensional despite the time-consuming simulation.
Chapter 3 Characterisation of non-torrefied and microwave torrefied biomass

This chapter presents the experimental procedure for microwave torrefaction process as well as an experimental method for calorific value determination and the proximate and ultimate analysis of the samples. The discussions on the comparison of physical and chemical properties for both samples are discussed.

3.1 Biomass sample

The biomass sample used in this study is white non-thermally treated wood biomass (Figure 3.1). The non-torrefied biomass was ground into a smaller size with an average size of 1.5 mm.

![Non-torrefied biomass sample](image)

Figure 3.1: Non-torrefied biomass sample

3.2 Torrefaction process

Based on the recent development in torrefaction technology as discussed in Section 2.2, the microwave torrefaction is considered as a promising technology with attractive benefits for the final product. Among the advantages of using the microwave as a heating source is the energy saving as it can be operated in shorter time to achieve the target temperature compared to the conventional heating. Despite those benefits, many researchers had reported that the product from the torrefaction process is a torrefied biomass that has intrinsic and chemical properties close to the low-grade coal. Thus, torrefied biomass may possess the similar tendency to self-heat comparable to coal. The possibility of the torrefied biomass to undergo self-heat had been discussed by Cruz Ceballos et al. (2015).
3.2.1 Equipment and experimental procedure for sample preparation

This research investigates the thermal behaviour of thermally treated biomass fuel using microwave torrefaction process in comparison to the non-torrefied material. The raw material used in this study was white non-thermally treated wood biomass. The woody biomass was ground into a smaller size with an average size of 1.5 mm. The raw material used as a received basis to prepare the microwave torrefied sample, where no pre-drying is necessary before torrefaction process. Research by Wang et al. (2012) verified that pre-drying is unnecessary for microwave torrefaction because the water content of the biomass fuel has no adverse effect on the torrefaction process.

Preparation of the torrefied sample was conducted at the Green Chemistry Centre of Excellence, Department of Chemistry, University of York. The sample was torrefied using Milestone ROTO Synth Rotative Solid Phase Microwave Reactor (Milestone Srl., Italy) as shown in Figure 3.2. The operating microwave power during the torrefaction process is 400 W. The high microwave power was chosen for this work to provide less torrefaction processing time. The process was done under vacuum condition at around 500 mbar at the highest heating temperature.

Figure 3.2: Microwave torrefaction reactor ROTO SYNTH (Milestone Srl., Italy)
The raw sample was weighed and placed in a 2 Litre glass flask. The glass flask was fitted with rotation device in the microwave chamber (as shown in Figure 3.3). The flask will rotate during the process to give a homogeneous heating to the sample. The reaction time was 6 minutes from room temperature to the final temperature (107°C). The by-products from the torrefaction were closely monitored during the process.

Figure 3.4: The liquid products were condensed in a water-cooled vacuum trap
At a temperature around 80 - 88°C, a clear liquid fraction was produced, and because of the increasing temperature, bio-oil was observed around 95-107°C. The liquid products were collected in water-cooled vacuum trap as shown in Figure 3.4. The process ended at this stage since this research focused on the microwave torrefied fuel generated at low temperature. The final weight of the torrefied biomass was recorded. The weight data was necessary to calculate the mass and energy yields as shown in Section 3.4.2.

3.2.2 Microwave torrefied biomass fuels

The change of colour for the microwave torrefied biomass sample is observed after the torrefaction process as shown in Figure 3.5. The colour of the microwave torrefied biomass fuel produced was darker compared to the non-torrefied sample. The change of the colour of torrefied biomass is because of the losses of the surface moisture, bound moisture, as well as light volatile gases during the torrefaction process (Nhuchhen, Basu, & Acharya, 2014). The torrefied sample produced for this study is brown with an inhomogeneous colour pattern.

![Microwave torrefied sample](image)

Figure 3.5: Microwave torrefied sample

This inhomogeneous colour of the torrefied sample is expected; due to the short reaction time for torrefaction process selected for this study. The observation had been explained in Satpathy et al. (2014) and Dhungana (2011), where the short reaction time, the non-homogenous nature of the biomass, as well as the selective heating mechanism of the microwave contributed to the inhomogeneous colour of
the torrefied sample. Also, the lower heating rate can also contribute to more homogenous products.

The varieties in the changes of colour for the torrefied product were reported as to be depending on the applied torrefaction conditions (Graham, 2015; Nhuchhen, Basu, & Acharya, 2014). This statement is supported by Cruz Ceballos et al. (2015); Shang et al. (2012); Phanphanich & Mani (2011) where they established that the colour of the materials that undergo torrefaction process were found to be darker with the increasing of torrefaction heating temperature. Besides that, the study by Satpathy et al. (2014) also showed the colour changes from dark brown to black when the microwave power level and the reaction time increased. Tumuluru et al. (2011) had identified the colour changes of raw material after the torrefaction process as a useful method to describe the degree of torrefaction, where the biomass turns from brown to black at 150°C – 300°C considering the chemical composition changes during the process.

It is worth mentioning that, a recent study by Sadaka et al. (2014), had published a detailed measurement of biomass colour changes, where they used biomass discoloration index to characterise the colour changes based on carbonisation temperature and reaction time during the carbonisation process. The index could potentially capture the extent of the biomass conversion through the degree of the colour changes at different carbonisation conditions. The use of the discoloration index can be extended on the biomass fuel produced using torrefaction process. Unfortunately, this index cannot be applied in this research because only one condition of torrefaction process was studied.
3.3 Methods to determine properties of biomass fuels

3.3.1 Calorific value

Calorific value or heating value is the amount of energy released during the combustion of material in a pure oxygen environment. The determination of calorific value was fulfilled based on ASTM D4809-13 using Isoperibol Oxygen Bomb Calorimeter (Parr Instrument Company, Parr 6200) that equipped with a microprocessor controller as shown in Figure 3.6 and Figure 3.7 shows the parts of bomb vessel.

The sample for the determination of calorific value was ground using a ball mill to about 125 µm. Sample with a weight of 1 gram was placed in a crucible. The crucible was put in the sample holder at attached to the bomb head. A 10 cm fuse wire was connected to the electrodes as an ignition source to start the combustion reaction of the sample in the vessel. Thus, the wire must touch the surface of the sample for ignition to occur. The bomb head was then carefully placed into the cylinder and sealed by hand tightening the screw-on cap. The bomb vessel was filled with pure oxygen at a pressure of 30 atm.

Figure 3.6: 6200 Isoperibol Bomb calorimeter (Parr Instrument Company, USA)
A water bucket was filled with 2 litres of water and placed into the bomb calorimeter. The bomb vessel was lowered into the bucket that filled together with the stirring mechanism. The stirrer will ensure a homogeneous temperature distribution of water in the bucket. Then the sample was ignited in pure oxygen within a sealed bomb vessel. The heat resulting from the ignition will cause a small increase in water temperature in the bucket that surrounding the bomb, which then recorded automatically by the sensor.

### 3.3.2 Proximate analysis

Proximate analysis is a method to characterise biomass by determination of moisture, fixed carbon, ash and volatile content in biomass sample. Proximate analysis determination was using thermogravimetric system following the standard method in ASTM Standard Test Method D5142. Based on the ASTM method, the moisture content, volatile matter, and fixed carbon can be determined successively in a single instrumental procedure. On the other hand, for fixed carbon, the value is a calculated from the difference between 100 and the total of the percentage of moisture, volatile matter and ash.
The analytical requirements for proximate testing using thermogravimetric had been summarised as the capability to record the weight of a sample as it is heated based on a designated temperature range and held isothermally at those temperatures, then change the sample’s environmental atmosphere from inert to oxidising (Cassel, Menard, & Earnest, 2012). Thus, in this study, TGA 4000 (PerkinElmer Inc, USA) is used to carry out the analysis as it meets the capability mentioned earlier.

Firstly, Nitrogen (99.5% purity) is used as the carrier gas to achieve the inert environment for the reaction. The nitrogen is purged at a rate of 20mL/min while the sample is heated from ambient temperature to $110^\circ$C. After reaching $110^\circ$C, the temperature was kept constant for 5 minutes; the weight loss seen here is due to the evaporation of moisture content in the sample. After 5 minutes, the sample was heated up to $900^\circ$C with a heating rate of $25^\circ$C/min in nitrogen. The condition was held for a minute where the complete release of volatile matter occurs at this stage. Then the supply gas was switched to Oxygen (99.5% purity) at 40mL/min, and the sample was heated up to $950^\circ$C at $10^\circ$C/min, to ensure the burn of the remaining fixed carbon. After that, the residual left from the process is known as incombustible ash.

### 3.3.3 Ultimate analysis

This analysis is performed to determine the elemental compositions by weight percentage of Carbon, Hydrogen, Nitrogen, Sulphur and Oxygen. It was carried out using Elemental analyser FLASH 2000 CHNS/O (Thermo Fisher Scientific Inc., USA). The basic premise behind the procedure is the combustion of the sample in a pure oxygen atmosphere, and the resultant gases are automatically measured (García et al., 2012).

Sample weighed 2-3mg was placed into the aluminium capsule and added with vanadium pentoxide ($V_2O_5$). Vanadium pentoxide acts as an oxidising agent to promote combustion process in the furnace. The analyser is equipped with an auto-sampler that will drop each sample sequentially by electronically controlled movements into a $900^\circ$C furnace. A small volume of oxygen is added to burn the sample that converts the sample into elemental gases. The element concentrations
were determined using a separation column and thermal conductivity detector equipped in the analyser.

### 3.3.4 True density and bulk density determination

The true density had been determined using AccuPyc II 1340 Automatic Gas Pycnometer (Micrometrics, Norcross, USA). The true density was determined by calculating the volume occupied by its mass without the volume of all pores. The volume is calculated using the pressure difference and the known volume of displacement.

The overall volume occupied by the mass of the sample, which included the pore and interstice volumes is the bulk density. The bulk density was determined by filling up the sample into a known cubic volume and measuring the amount of the samples without the tamping the samples.

### 3.4 Results and discussion

#### 3.4.1 Calorific value

Figure 3.8 shows the results from the experiment using bomb calorimeter the increase in the calorific value of the microwave torrefied sample. The calorific value of the non-torrefied sample is 18.4MJ/kg and 22.0MJ/kg for torrefied biomass. The calorific value of the microwave torrefied biomass had increased to around 20% compared to its raw material.

![Figure 3.8: Calorific value of the samples](image)

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The calorific value determined for the torrefied sample in this study is in the range of values reported by several researchers such as Ren et al. (2013) for torrefied Douglas fir pellet using microwave reactor; Kopczyński et al. (2015) for torrefied willow using fixed bed reactor; Wang et al. (2013) for torrefied sawdust in an oxidative torrefaction process and Jones et al. (2012) for torrefied willow using reactor tube for conventional torrefaction. Even though the material and method of torrefaction are different for the mentioned researchers, the values reported were in the range as stated by Kleinschmidt (2011) in his review, which is between 20-24MJ/kg.

The increased of calorific value had been described by Eseyin et al. (2015) and Wang et al. (2013) was due to the removal hemicellulose content in biomass, where hemicelluloses go through most reactive devolatilization and carbonisation under 250°C. The calorific value values increased linearly with the torrefaction temperature in the laboratory scale study of conventional torrefaction of reed canary grass, wheat straw and willow by Bridgeman et al. (2008). This pattern is expected as the moisture contents of the samples also decrease with the increased of the torrefaction temperature. The same relationship was observed in the recent study by Kopczyński et al. (2015) on torrefied willow at a different temperature where the calorific value of 19.33MJ/kg for torrefaction temperature of 200°C was observed and increased to 26.5MJ/kg when applied with torrefaction temperature of 300°C.

However, it is a different case for the application of microwave induced pyrolysis on rice straw at 400-500W microwave power reported by Huang et al. (2008). They indicated that there is no correlation between the calorific value and the microwave power/temperature. The reduction of calorific value under the torrefaction condition of 400-500W microwave power can be seen in their findings. They had concluded that the findings might suggest that when microwave power is higher than 400W, some of the fixed carbon content can also be pyrolyzed, which lead to the reduction in the calorific value values. Moreover, their research showed that the water content of biomass fuel used did not have a substantial effect on the calorific value. On the other hand, Wang et al. (2012) observed the increase of the calorific value of the biomass products due to water content at lower microwave power levels (below 300W)
A study done by Lanigan (2010) had shown the efficiency of using microwave irradiation pyrolysis compared to conventional pyrolysis. Based on the research, the similar calorific value could be produced at temperatures 100°C lower than those produced using conventional pyrolysis. For conventional pyrolysis, the calorific value increases with temperature and residence time (Arias et al., 2008). Besides that, the results of the microwave torrefied biomass determined in this work is comparable to calorific values determined by low-temperature pyrolysis using pine wood and coconut fibre in (Liu & Han, 2015). The present result shows that the low-temperature microwave torrefaction has potential to produce solid biofuel similar to the one using other types of thermal pre-treatment.

### 3.4.2 Mass and energy yield

The torrefaction process had been proved to reduce the mass and energy yield of the final product. There are several correlations between the microwave torrefaction conditions and the percentage of mass and energy yield of biomass torrefied material. For example; the percentage of mass yield reduced with the increased of the torrefaction temperature in research by Poudel & Oh (2014) as well as van der Stelt et al. (2011). In the other hand, Satpathy et al. (2014) found the mass yield reduced when the reaction time increased. In addition to the torrefaction conditions, findings in Satpathy et al. (2014) presented that the moisture content also has a significant influence on the pattern of mass yield in torrefaction process, where most of the higher mass yield achieved at higher moisture content.

Mass and energy yield shows in Figure 3.9 was calculated for the microwave torrefaction process using the Eq. 3.1 and Eq. 3.2 based on Bridgeman et al. (2008).

\[
Mass \ yield \ (M) = \frac{\text{Mass after torrefaction}}{\text{Mass of raw sample}} \times 100 \quad (\text{Eq. 3.1})
\]

\[
Energy \ yield \ (E) = Mass \ yield \ (M) \times \frac{HHV \ after \ torrefaction}{HHV \ of \ raw \ sample} \times 100 \quad \text{(Eq.3.2)}
\]
Chapter 3: Characterisation of non-torrefied and microwave torrefied biomass

Mass balance calculation shows that 73% mass and 87% of energy retained in solid form, while 27% mass and 13% of energy converted to liquid and gases products. The mass and energy yield calculated in this research is in agreement with the microwave torrefaction research of rice straw and pennisetum by Huang et al. (2012) and Ren et al. (2012a). In addition to that, Ren et al. (2014) also reported the energy yield of microwave torrefied corn stove ranged between 87.03 to 97.87% subjected to the torrefaction holding time and temperature chosen. They also observed the higher energy yield of the torrefied biomass conducted in lower temperature reaction. Besides that, the mass yield of torrefied wood showed a significant reduction when higher torrefaction temperature applied (Ben & Ragauskas, 2012).

Based on the findings by Huang et al. (2012) the optimum condition for needed microwave power level to produce 70% of the mass is just only 150W with 10 minutes reaction time. Thus, their finding correlated with the mass yield determine in this study, with the processing time just 6 minutes using higher microwave power, which is 400W. Thus, the mass and energy yields from this study provide ample support to validate the findings that torrefaction process able to retain the energy in solid form. However, the percentages of the mass and energy yield are widely influenced by the parameters of torrefaction process such as temperature and reaction time.

Figure 3.9: Mass (M) and energy (E) balance from the torrefaction process
3.4.3 Proximate analysis

3.4.3.1 Discussion on TG-DTG curves for proximate analysis

Figure 3.10 shows the temperature profile during the proximate analysis based on the procedure explained in Section 4.2. It can be observed from the percentage of weight loss curve that the temperature profile of the microwave torrefied sample shifted to a higher temperature. Figure 3.10 (a) shows that TG profiles are divided into three noticeable phases, correspond to moisture loss (30ºC - 110ºC), active (110ºC - 400ºC) and lastly the passive pyrolysis stage.

The first phase is from room temperature to 110ºC to indicate water evaporation phase. During this phase, the sample underwent the extraction of moisture and adsorbed water of the sample. It is shown in Figure 3.10 (a) that in the first phase, the mass loss of non-torrefied sample is more than the microwave torrefied sample due to the torrefaction process that drives out the moisture. At around 250ºC, the percentage of weight loss for both samples is slightly similar, and after that, the active thermal decomposition took place after that.

The different of weight losses for both samples is due to the drying process and the removal of low-molecular-weight volatiles during torrefaction (Wang et al., 2012). Based on Figure 3.10 (b), the maximum peak of for non-torrefied sample is 370ºC, whereas it is 366ºC for microwave torrefied sample. The rate of weight loss reaches a maximum as the rapid volatilization occurred, accompanied by the formation of the carbonaceous residue (Kopczyński et al. 2015).

In the active pyrolysis phase, the main decomposition reactions occurred and most of the organic compound related to hemicellulose and cellulose decomposed. The higher mass loss in the decomposition of the non-torrefied sample is expected as it contains higher volatile matter (84.6%) compared to the torrefied sample (68.7%). In the last phase, the passive pyrolysis can be monitored as there is no significant mass loss and the residues are decomposed to the final temperature. At this higher temperature, the complete decomposition of lignin content occurred and followed by the oxidation of the char formed by pyrolysis process (Saldarriaga et al., 2012).
Based on the temperature profile in Figure 3.10 (a), after heating to 800°C approximately 30% of microwave torrefied sample remained while only approximately 15% of the non-torrefied sample was left. Therefore, based on the TGA results, the non-torrefied sample is more reactive to thermal process compared to the microwave torrefied sample. The increase of reactivity contributed to a more stable torrefied material after the torrefaction process.

3.4.3.2 Discussion on the composition of the samples

The results of the proximate analysis of both samples were shown in Table 3.1, and was compared to the range values reported by Vassilev et al. (2015). The properties of the torrefied biomass found to be in between wood and coal except for volatile matters, which is relatively higher than the value of coal. However, the upgraded properties compared to wood are a desirable characteristic needed for the usage of torrefied biomass as alternative fuel.

The moisture content of the non-torrefied sample is 6.9% and decreases to 2.2% after the torrefaction process. The reduction is more than 50% compared to the non-torrefied sample. The reduction is expected due to one of the advantages of torrefaction process that include reduced moisture content. Besides that, torrefaction is also proven to reduce the water adsorption capacity and acquired hydrophobicity nature of the resultant products (Satpathy et al. 2014; Huang & Hsu 2012; Li et al. 2012; van der Stelt et al. 2011). In addition to that, Tumuluru et al. (2011) suggested that hydrophobicity nature was due to the loss of the low-volatility components and extractive during the torrefaction.
Table 3.1: Proximate analysis of non-torrefied and microwave torrefied samples in comparison with coal

<table>
<thead>
<tr>
<th>Proximate Analysis (wt.%)</th>
<th>Non-torrefied sample</th>
<th>Microwave torrefied sample</th>
<th>Coal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture content</td>
<td>6.90</td>
<td>2.20</td>
<td>0.4 – 20.2</td>
</tr>
<tr>
<td>Volatile matter</td>
<td>84.60</td>
<td>68.70</td>
<td>12.2 – 44.5</td>
</tr>
<tr>
<td>Fixed carbon</td>
<td>3.90</td>
<td>18.60</td>
<td>17.9 – 70.4</td>
</tr>
<tr>
<td>Ash*</td>
<td>4.70</td>
<td>10.5</td>
<td>5.0 – 48.9</td>
</tr>
</tbody>
</table>

* Content is obtained by difference

¹Source: Vassilev et al. (2015)

Volatile matters refer as the components released, including combustible gases (C₅H₇ gases, CO or H₂) and some incombustible part (CO₂, SO₂ or NOₓ) when the sample heated at high temperature. The volatile matters in the torrefied sample are slightly lower than the non-torrefied sample, due the changes occur through product’s devolatilization during the torrefaction process. During the microwave torrefaction process, the condensable volatile matters were condensed into a liquid such as bio-oil. The values of the volatile matter in the both samples are higher than the one found in coal. According to Chiang et al. (2012), the material with higher volatile matter will has lower ignition temperature and high ignition stability to produce synthesis gas during the biomass gasification process. They also added that, the finding indicated that the coals exhibit less volatility during thermal conversion process.

The composition of volatile matters observed to reduce from 84.6% to 68.7%, which is a reduction of 18.79% for the microwave torrefied sample compared to the non-torrefied sample. The reduction was due to the decomposition of the hemicelluloses to volatiles matter while partly decomposition occurred for cellulose and lignin. However, the degree of decomposition is subject to the parameters of torrefaction process such as reaction temperature (Huang et al., 2012). The study by Wang et al. (2012) indicated that with increased processing time and certain power level; the reaction could become more complete, which resulted in a further degree of decomposition of lignocelluloses that eventually lead to more removal of volatiles.
from the biomass. The reduction of the volatile matters in the microwave torrefied biomass may suggest the lower reactivity of the material.

Fixed carbon is the carbon that is left after volatile materials are driven off during the torrefaction process due to the conversion of hemicellulose into the more thermally stable compound. In this work, the fixed carbon content in the torrefied sample showed a significant increase compared to its non-torrefied sample. The fixed carbon in the microwave torrefied sample is almost five times higher than the non-torrefied sample. Based on the proximate analysis done, the fixed carbon in the non-torrefied sample is 3.9% whereas for the microwave torrefied sample is 18.6%. The fixed carbon found in microwave torrefied sample was within the range of the one found in coal. This showed that thermal properties of the material had been upgraded comparable to coal.

Ash is an inorganic part of the sample, which left after the complete combustion occurs. Ash content in the torrefied sample is over two times higher than to the non-torrefied sample. Where the ash content in torrefied biomass samples is 10.5% compared to only 4.7% in the non-torrefied sample. The increased value of the ash content can be seen in most of the torrefied biomass either using microwave induced method or conventional torrefaction process (Poudel & Oh, 2014; Satpathy et al., 2014; Huang et al., 2012; Fisher et al., 2012). Furthermore, Wu et al. (2015) stated that ash content of microwave pyrolysis biomass at is higher than the one produced using conventional pyrolysis process. Thus, it can be suggested that the method of heating using microwave have some effect on the ash content.

Based on the research of self-heating on coal; higher ash content will lead to slower self-heating rate (Beamish, Lin, & Beamish, 2012) as the ash can block the active site. Thus the ash content reduced the reactivity of the torrefied biomass as the accumulation of the ash on the surface reduces the direct contact for the oxidation process. However, high ash content will also lead to several problems such as a reduction in heating value, and the ash deposits can cause extensive equipment maintenance during the thermal treatment (García et al., 2012). Consequently, Khan et al. (2009) had concluded that the difficulty related to high ash content in biomass as the biggest technical challenge.
Thus, it is agreeable to findings by earlier researchers that acknowledged the torrefaction process as a highly proven method to reduce moisture content and volatile matter as well as increase the fixed carbon and ash content (Cruz Ceballos, Hawboldt, & Hellleur, 2015; Kopczyński, Plis, & Zuwala, 2015; Peng et al., 2013; Agar & Wihersaari, 2012; Bridgeman et al., 2008).

### 3.4.4 Ultimate analysis

Table 3.2 shows the comparison of the chemical composition of the samples as compared to the range values reported for coal in Vassilev et al. (2015). Sulphur content was below the detected limit of the equipment. Therefore, the sulphur content is discarded in the analysis. Based on a review by Vassilev et al. (2015) and Yin et al. (2008), the low Sulphur content is an excellent characteristic in biomass fuel as it leads to lower SO$_2$ emissions and limited production of fine particulates, deposit formation and corrosion during the thermochemical process. Therefore, it is safe to consider that the microwave torrefied sample in this study demonstrates a desirable characteristic as alternative fuels to coal.

During torrefaction process, the biomass loses oxygen and hydrogen due dehydration of water and organic reaction products, for example, acetic acid, furans, methanol and gases that contain a substantial amount of oxygen such as CO$_2$ and CO (Bergman & Kiel, 2005). The percentage of H and O content in the microwave torrefied sample is less than the non-torrefied sample. While the nitrogen and carbon content are higher in the microwave torrefied sample compared to the non-torrefied sample.

The trend presented in Table 3.2 are similar to the findings by Prins, (2005); Ren et al. (2012); Huang & Hsu (2012); Poudel & Oh (2014); Cruz Ceballos et al. (2015); Kopczyński et al. (2015) where similar pattern of increased carbon and decreased of oxygen content in torrefied biomass has been found; despite the method of torrefaction used.
Table 3.2: Ultimate analysis of non-torrefied and microwave torrefied samples in comparison to coal

<table>
<thead>
<tr>
<th>Ultimate Analysis (wt.%)</th>
<th>Non-torrefied sample</th>
<th>Microwave torrefied sample</th>
<th>(^{1})Coal</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>54.4</td>
<td>61.50</td>
<td>62.9 – 86.9</td>
</tr>
<tr>
<td>H</td>
<td>7.70</td>
<td>6.80</td>
<td>4.4 – 29.9</td>
</tr>
<tr>
<td>N</td>
<td>1.70</td>
<td>3.80</td>
<td>3.5 – 6.3</td>
</tr>
<tr>
<td>O(^{*})</td>
<td>36.2</td>
<td>27.90</td>
<td>0.5 – 2.9</td>
</tr>
<tr>
<td>O/C</td>
<td>0.67</td>
<td>0.45</td>
<td>-</td>
</tr>
<tr>
<td>H/C</td>
<td>0.14</td>
<td>0.11</td>
<td>-</td>
</tr>
</tbody>
</table>

\(^{*}\) Content is obtained by difference

\(^{1}\)Source: Vassilev et al. (2015)

The carbon content of the microwave torrefied sample showed an increase of roughly 13% compared to the non-torrefied sample. The increase is due to the carbon preservation after torrefaction process completed. Besides that, nitrogen content rose from 1.7% in the non-torrefied sample to 3.8% in microwave torrefied sample.

According to ultimate analysis of the torrefied biomass by (Prins, 2005; Kopczyński et al. 2015), they had found a relationship between the torrefaction temperature and the amount of oxygen and carbon left in the torrefied biomass. Where the higher the torrefaction temperature is applied, the lower the oxygen content left in the sample. While for the carbon content, it increased with the increase of torrefaction temperature.

The findings in this research followed the pattern in Wang et al. (2013) where the hydrogen and oxygen content in torrefied sample decreased, and this may be due to the release of volatiles that rich in hydrogen and oxygen such as water and carbon dioxide. The release of water can be proved based on the earlier discussed findings in the proximate analysis that showed a reduction of 68% moisture content in the torrefied sample compared to its raw material.

Moreover, the common trend in torrefied sample is a reduction of H/C and O/C ratio compared to its raw material can be observed in several studies on torrefied biomass.
such as (Satpathy et al., 2014; W.H. Chen et al., 2013; Arias et al., 2008; Huang et al., 2008; Prins, 2005). The reduction of hydrogen and oxygen had been described due to the water, carbon dioxide and carbon monoxide production during the torrefaction process.

The H/C ratio for the non-torrefied sample is 0.14, and it decreased to 0.11 in the torrefied material. The reduction of 21.43% H/C ratio is due to the removal of moisture content during torrefaction where decomposition of hemicellulose and dehydration of lignin and cellulose simultaneously occurred. The removal of the moisture content and the reduction of the H/C ratio leads to higher calorific value. Therefore based on a study by Poudel & Oh (2014) the O/C and H/C ratios are found to reduce with an increase of the torrefaction temperature where further removal of hydrogen and oxygen occur at a higher temperature. According to Tumuluru et al. (2011) the reduction of O/C and H/C also reduced the energy loss, produced less smoke and water vapour during combustion and gasification process. Therefore, those reductions are desirable characteristics in the utilisation of biomass fuels as a substitute for coal.

The reduction of O/C ratio in microwave torrefied sample increases the resistance to thermal degradation. Therefore, one of the objectives of torrefaction processes is to produce biomass fuels that resistance to thermal degradation. The O/C ratio in the torrefied sample is 0.45 compared to 0.67 for its non-torrefied sample. The lower O/C ratio contributes to the higher yield during gasification compared to its raw material (Prins et al., 2006a). The work by Satpathy et al. (2014) presented a relationship between H/C and O/C ratios, where the ratio decreased with increase in power and reaction time of microwave torrefaction process.

The ultimate analysis shows that the torrefaction process had improved the quality of the biomass fuels. According to Valix et al. (2016) the H/C and O/C ratios of various types of coal are between 1.02 - 0.078 and 0.4 - 0.08 respectively. The value of the H/C and O/C ratios for the microwave torrefied lies in the ranges stated, which reflects that its properties had improved closer to coal.
In addition to that, similar to the finding by Valix et al., (2016) that indicated the reduction of the H/C content suggested the increase in aromaticity which leads to the better resistance to biological degradation. The same findings of increases in the degree of aromaticity for the sludge that had to undergo the pyrolysis process can be found in (Kan, Strezov, & Evans, 2016). A recent study of the off-gas emissions during storage of torrefied biomass had proved that the biological degradation was not the reason that contributed to the emission (Tumuluru et al., 2015). Thus, it can be concluded that the microwave torrefaction process had improved the resistance toward biological degradation.

In term of a tendency to the self-heating, García Torrent et al. (2016) have observed that biomass with higher H/C has higher tendency to self-ignite and this finding seem to contradict to the behaviour witnessed in coal. However, they suggested that the inconsistency could be due to different mechanisms of self-heating development that lead to self-ignition in biomass in comparison to coal. Thus, according to that observation, the non-torrefied sample will have a higher tendency to self-heat compared to microwave torrefied sample. However, this initial assumption needs to be clarified using thermal analysis to get an accurate measurement of the tendency of the samples to self-heat.

3.4.5 True density and bulk density

Based on the experiments done, the true density of the non-torrefied sample is 1451 kg/m$^3$ and 1442 kg/m$^3$ for microwave torrefied sample. It can be seen here that the true density of the microwaved torrefied sample is lower than the non-torrefied sample. The reduction is expected due to the lower volatile matters in the microwave torrefied sample, which resulted in more pore volumes.

The bulk densities calculated are 443 kg/m$^3$ and 500 kg/m$^3$ for non-torrefied sample and microwave torrefied sample respectively. The increased of the bulk density proved that the energy density increased after torrefaction process (Satpathy et al., 2014; Medic, 2012; Shang, 2012). The reduction of the bulk density due to the evaporating water and volatiles matters in the biomass by the torrefaction process.
3.5 Summary

- The torrefaction process upgrades the quality of the biomass, where the calorific value of the microwave torrefied sample increases to 22 MJ/kg from the 18.4 MJ/kg in the non-torrefied sample. The bulk densities are 443 kg/m³ and 500 kg/m³ for non-torrefied sample and microwave torrefied sample respectively.
- The upgraded biomass fuels produced from the torrefaction process is more reactive compared to the non-torrefied fuels. The TG curve of the microwave torrefied sample shifted to higher temperature region compared to non-torrefied sample, which indicated the higher reactivity of the sample non-torrefied sample.
- The moisture content reduced from 6.9% in the non-torrefied sample to 2.2% of microwave torrefied sample. Volatile matters reduced from 84.6% in the non-torrefied sample to 68.7% in microwave torrefied sample.
- The fixed carbon content in the non-torrefied sample is 3.9% whereas, for the microwave torrefied sample the content increased to 18.6%.
- The ash content in torrefied biomass samples is 10.5% compared to only 4.7% in the non-torrefied sample.
- O/C and H/C ratio decreased after the torrefaction process, which is one of the desirable characteristics of using torrefied biomass fuel as an alternative to coal. The O/C ratio is 0.67 in the non-torrefied sample and 0.45 in microwave torrefied sample. Whereas, the H/C ratio is 0.14 in the non-torrefied sample and 0.11 in microwave torrefied sample.
- The true density of the non-torrefied sample is 1451 kg/m³ and 1442 kg/m³ for microwave torrefied sample. Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis
Chapter 4 Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis

This chapter consists of the evaluation of the samples reactivity based on their thermal behaviour using thermogravimetric (TG) analysis and the kinetic parameters determination of the thermal decomposition of biomass in the air and inert condition using nitrogen as carrier gas. The objectives of this chapter are:

(a) to evaluate the thermal decomposition behaviour of the torrefied and non-torrefied biomass sample in air and nitrogen;

(b) to determine the characteristic temperature of both samples to analyse their reactivity; and

(c) to establish the kinetic parameters of the torrefied and non-torrefied biomass sample to evaluate their propensity of self-heating.

4.1 Equipment and method used for thermal behaviour analysis

TG analysis is used to obtain the critical parameters based on their heating behaviour. The thermogravimetric analysis of the sample was performed using a TGA 4000 (Perkin Elmer, USA). The weight loss of sample due to thermal degradation in the different carrier gas at the various heating rate was observed in this experiment. Only a small amount of sample needed for each test as this will ensure the uniformity of the sample. Samples weighing around 18-20mg were used for throughout the analysis.

There are three methods conducted in this study to analyse the thermal behaviour of the samples. The methods employed are as below:

a) Thermal behaviour in air and inert condition

This approach was used to assess the reactivity of the material that prone to self-heat. The reactivity of the samples is evaluated by identifying the mass loss behaviour of the samples when heated at fixed heating rate using air and nitrogen as a carrier gas. Nitrogen is used as a comparison
of the thermal behaviour in an inert environment. The samples were subjected to several heating rates namely 1°C/min and 5°C/min from ambient temperature to 900°C in the air. The flow of carrier gas is fixed at 50ml/min throughout the experiment. The heating rate of 1°C/min was chosen to represent the slow heating rate of self-heating behaviour, which allows a fully developed exothermic reaction in order to determine the temperature of initial combustion (TIC). While 5°C/min was chosen for the evaluation of the thermal characteristic of both sample in air and inert condition.

b) **Susceptibility evaluation in oxygen**

The TG analysis in the oxygen was done to fine a single point oxidation temperature. The carrier gas used in this analysis is oxygen, and the flow is fixed at 50ml/min. The heating rate is set to 40°C/min. The same condition applied to both samples.

c) **Kinetic parameter determination**

Samples were heated from ambient temperature to 900°C in air at a heating rate of 5, 10, 20 and 30°C/min. The flow of carrier gas is fixed at 50ml/min throughout the experiment. The selected heating rates are chosen based on the study done by Benitez, (2014) where the heating rate higher than 50°C/min had been proved to produce TGA curves with high variability and rapid changes.

Each experimental run was repeated at least three times, and the curves for each heating rates were compared to check the reproducibility. The reproducibility was verified by comparing the temperature, the conversion and weight derivative at peak DTG curve for each heating rate. In addition to that, a constant gas flow is chosen to achieve a low noise TG reading (Mansaray & Ghaly, 1999).
4.2 Evaluation of thermal reactivity

4.2.1 Thermal decomposition characteristic

The method employed for the analysis of intrinsic reactivity was explained in Section 4.1. Air was used as carrier gas because this study focuses on the behaviour of the sample during storage in an air atmosphere. Nevertheless, the analysis was extended using nitrogen to compare the thermal decomposition behaviour of the sample in an inert environment.

4.2.1.1 Thermal decomposition in air

For the evaluation purposes, the temperature profiles of thermal decomposition in the air at 5°C/min is presented to evaluate the reactivity of the samples in air at a low heating rate. The low heating rate for the reactivity evaluation in the air had been chosen to allow a fully developed exothermic reaction as suggested by Della Zassa et al. (2013).

Figure 4.1 shows the temperature profile of both sample during TG analysis in the air at 5°C/min. The graph indicates that the thermal decomposition of microwave torrefied sample shifted to a higher degree compared to the non-torrefied sample. This pattern represents the sample’s reactivity in an air stream at 5°C/min, where non-torrefied sample seems to decompose in the air at a lower temperature than the microwave torrefied sample. Based on the TG analysis, 10% weight loss was identified at the temperature of 252°C for the non-torrefied sample. However, for the microwave torrefied sample, the same weight loss was achieved at 267°C. Therefore, based on the TG analysis in air, the non-torrefied sample is more reactive in the air compared to the microwave torrefied sample.

The same pattern was reported by several researchers when comparing the temperature profile of the torrefied sample to its raw materials such as for Cruz Ceballos et al. (2015); Kopczyński et al. (2015); Mašek et al. (2013); Ren et al. (2013). A study by Ren et al., (2013) indicates the relationship of the degree of the shift with the severity of torrefaction condition and the same relationship can be observed in Cruz Ceballos et al. (2015). In both studies, the decomposition of the
torrefied biomass seems to shift to a higher degree for the biomass that exposed to higher reaction temperature during the torrefaction process.

From the rate of mass loss shown in DTG graph (Figure 4.2), three clear peaks can be observed for both samples. The first peak represents the moisture driven out from both samples. Based on the graph, more derivative weight loss occurred in the non-torrefied sample at the first peak compared to the microwave torrefied sample because of the different value of moisture content. Based on the proximate analysis, the moisture content of the non-torrefied sample is 6.9%, while for microwave torrefied sample only 2.2%. Thus, increasing the temperature from ambient to about 110°C resulted in the weight losses due to the moisture content in the samples.

Figure 4.1: Temperature profile for thermal decomposition in air at 5°C/min

Figure 4.2: Derivative weight loss in air at 5°C/min
Chapter 4: Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis

The second and third peaks represent the thermal decomposition of biomass composition that decomposed at different temperatures. The hemicellulose is decomposed at a lower temperature between of 225 – 325°C, cellulose at 305 – 375°C and lignin steadily decomposed over the temperature range of 250 – 500°C (Prins et al., 2006b). The main decomposition region for both samples is between the ranges at approximately 240°C to 380°C, which the second peak fall. Hemicellulose decomposed during this main decomposition stage. The third peak represents the cellulose decomposition that occurred at a higher temperature range. In addition to that, lignin decomposed gradually during the thermal analysis contributed to the noticeable derivative weight loss at both peaks.

The maximum temperature for weight loss based on results presented in Figure 4.2 shows a shift of maximum weight loss of 337°C for the non-torrefied sample to 346°C for microwave torrefied sample. Thus, it indicates that a higher temperature needed by microwave torrefied sample to undergo the highest yield of volatile matters during the thermal decomposition in air. The higher temperature needed by microwave torrefied was considered to be contributed by the volatile matters driven out during the torrefaction process.

4.2.1.2 Thermal decomposition in nitrogen

Based on the analysis of both samples under an inert environment, the percentage weight loss is shown in Figure 4.3 and percentage of derivative weight loss is presented in Figure 4.4. The thermal decomposition in nitrogen follows the patterns of temperature profiles in biomass pyrolysis. The same pattern of temperature profile shifted to a higher degree for microwave torrefied sample can be seen in Figure 4.3. This finding is similar to the analysis done in the air as shown earlier in Figure 4.1. However, in a nitrogen atmosphere, we can see that both samples did not achieve full weight at the final temperature. The solid residue yields are about 15% for non-torrefied sample and 25% for microwave torrefied sample.
Figure 4.3: Temperature profile for thermal decomposition in nitrogen at 5°C/min

Three regions can be witnessed in Figure 4.4. Those regions correspondent to the moisture driven out process, active and passive pyrolysis. The first region follows the same pattern as thermal decomposition in air. The active pyrolysis phase occurs at the second region which lies in a range from 120°C to 425°C for heating rate 5°C/min. In this region, only one clear peak is observed, which represents the highest derivative weight loss. Furthermore, there is no third peak can be seen in DTG curve of the thermal decomposition in nitrogen due to the reaction in air added the stage of charring (Liu et al., 2016).

Similar to the thermal decomposition in the air, the temperature where the highest derivative weight loss occurs shifted to a higher temperature for the microwave torrefied sample. Based on Figure 4.4, the temperature is 370°C for non-torrefied sample and 372°C for microwave torrefied sample. The value of the temperature that higher weight loss occurred for both sample in nitrogen is slightly higher compared to the analysis done in air. The pattern found here proved that higher reaction temperature is needed to achieve the highest derivative weight loss in an inert environment.
Based on the analysis of temperature profiles presented here, the non-torrefied sample can be proposed as a higher reactivity material compared to the microwave torrefied sample. Thus it can be said that torrefaction process helps to upgrade the thermal properties of the microwave torrefied sample, which leads to a higher thermal stability due to less volatile matters (Agar & Wihersaari, 2012).

### 4.2.2 Temperature of initial combustion (TIC)

The temperature of initial combustion is essential in reactivity analysis because it reflects the starting point of the volatile matters to decompose after moisture driven out from the material. The more reactive material will have lower TIC, due to the ease of the material to react in the air. Determination of the TIC was done based on the thermal degradation behaviour of the samples at 1°C/min in air. The temperature profile used to find the TIC is shown in Figure 4.5.

Following the method to determine the TIC as explained in Ramírez et al. (2010), TIC for the non-torrefied sample is 268°C, whereas 283°C for the microwave torrefied sample. These values agree with the analysis done in 5°C/min, which concluded that the non-torrefied sample is more reactive compared to the non-torrefied sample.
Chapter 4: Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis

Figure 4.5: Determination of temperature of initial combustion (TIC)

The increase in the initial combustion (TIC) for the microwave torrefied sample is consistent with the observed reduction of the volatile matter as been shown in Table 3.1. The increase of temperature had suggested that the decrease of the combustible volatile matters due to the torrefaction process. The same behaviour can be found in Liu et al. (2016) where the temperature of initial combustion of the torrefied biomass also shifted to a higher temperature in their research. They also mentioned that the finding might result in higher combustion efficiency and reduction of emissions in the combustion process of the torrefied biomass.

4.2.3 Temperature at maximum weight loss (TMWL)

To compare the propensity of self-heating for the non-torrefied sample and microwave torrefied sample. The TG analysis in the air is done at 1°C/min based on method explained in Section 4.1(a). The heating rate of 1°C/min is chosen to portrait a prolonged heating in an air atmosphere. The slow heating rate allows a more rapid thermal energy supplied to the sample during the decomposition process (Slopiecka, Bartocci, & Fantozzi, 2012) in longer reaction time.
Figure 4.6: Determination of temperature of maximum weight loss in air at 1°C/min

Figure 4.6 shows the DTG analysis at 1°C, where the maximum derivative weight loss is 310°C for non-torrefied sample and 312°C for microwave torrefied sample. Thus, the temperatures of maximum derivative weight loss demonstrate that microwave torrefied sample is less reactive compared to the non-torrefied sample. The temperature difference is too small, only exceed 2°C for microwave torrefied sample compared to the non-torrefied sample. This slight difference makes it hard to distinguish the reactivity between the two samples. The differences suggested that both materials reacted slightly similar in the air. The samples need to be tested under oxygen stream in order to get a clear justification on this matter which is discussed in Section 4.2.4. However, the TMWL determined for both samples from the TG analysis are in agreement with the finding by DTI (2013) for pine chips torrefaction that showed sign of self-ignition at temperatures above 200 °C.

### 4.2.4 Characteristic oxidation temperature (T\textsubscript{charac})

To get a single point oxidation temperature, the TG analysis using oxygen as a carrier gas is done. The temperature profile of the analysis is shown in Figure 4.8. The characteristic oxidation temperature is an important parameter since it can be used with the activation energy to give a comparative assessment of the self-heating propensity (Saddawi et al., 2013).
Based on the graph presented in Figure 4.7, a single oxidation temperature can be assigned to the sample. $T_{\text{char}}$ for the non-torrefied sample and microwave torrefied sample are 331°C and 329°C respectively. The results of $T_{\text{char}}$ for the samples imply that the microwave torrefied sample is more reactive compared to the non-torrefied sample. The finding, similar to the study by Cruz Ceballos et al. (2015), where they found that torrefaction had caused depletion of volatiles component which increases the available surface of active sites for oxidation which will lead to self-heating. The occurrence can only be seen clearly when the TG analysis was done in oxygen stream.

![Figure 4.7: Determination of $T_{\text{char}}$ under oxygen stream at 40°C/min.](image)

However, the reactivity susceptibility of the material in the air based on $T_{\text{char}}$ contradict with the earlier conclusion based on TIC and TMWL that reveals non-torrefied sample is more reactive than the microwave torrefied sample. Therefore, further investigation using kinetic parameters analysis is needed to confirm the reactivity of the samples in air.
4.3 Results of kinetic parameters using thermogravimetric analysis

4.3.1 Background on determination of kinetic parameters

It is important to understand the reaction kinetic of the oxidation process that takes place at a low temperature in order to assess the risk of self-heating events that lead self-heating. Based on the study of intrinsic reactivity of coal chars in Sima-Ella et al. (2005), the activation energy affected the temperature sensitivity of the material, while the pre-exponential factor is related to the material structure (Sima-Ella, Yuan, & Mays, 2005). Consequently, due to the similarity of the properties of the microwave torrefied sample and coal, it is suggested that its behaviour will follow the same intrinsic reactivity. Therefore, in this study, it is sufficient to characterise the reactivity by its activation energy value only for both samples. However, due to the limited volume of samples, the kinetic study using adiabatic oven test as described in BS 15188 (2007) cannot be applied.

Thermogravimetric (TG) analysis is widely used to obtain kinetic parameters for the low-temperature oxidation. The mass changes of the material obtained from TG analysis at several heating rates is seen as a useful tool for determination of kinetic parameters of solid-state reactions (Carrier et al., 2016; Prins, 2005). There are many different methods developed to quantify the mass changes reactions and commonly categorised as model-fitting and model-free methods. For the model-fitting method, different models are fits to the experimental data and the model giving the best statistical fit is selected. The evaluations of the model were done from which the activation energy ($E_a$) and frequency factor (A).

The model fitting methods were broadly used because kinetic parameters can be calculated directly from the thermogravimetric analysis results. However, these methods have several disadvantages. One of the major advantages, has been discussed in Heydari et al. (2015) is their inability to select the appropriate reaction model uniquely, and the comparison of results from these models can be challenging as the wide range of kinetic parameters in non-isothermal data.
Hence, in this research, a model-free method is adopted to calculate the kinetic parameter of the sample decomposition. This model is much simple to apply and can allow the determination of the kinetic parameters for different constant extents of the conversion without considering any particular form of reaction model (Heydari, Rahman, & Gupta, 2015). The determination is possible because this method involves reaction rate that depends on the extents of conversion to product, where several kinetic curves involve in the calculation of the activation energy at the different heating rates on the same value of the conversion. The method of using multiple heating rates is proven more efficient compared to single heating rate.

Thus for non-isothermal experiments, each run need to be done under the same experimental conditions, such as same sample weight, purge gas rate and sample size, then the only variable is the heating rate. The temperature sensitivity of the reaction rate depends on the extent of conversion to products. A parameter called conversion degree (\(\alpha\)) is defined for the subsequent analysis of each phase of the decomposition process.

Changes of air to the inert environment resulted in the increase in values of activation energy, as the decomposition in the air is significantly faster compared to the decomposition process in an inert atmosphere. Thus, in this section the discussion of the comparison of the behaviour of the sample for the thermal decomposition in inert and air atmosphere. For the numerical modelling of the self-heating behaviour, the kinetic parameters for the thermal decomposition in air atmosphere are used.

### 4.3.2 Kinetic analysis method

The simplest way for modelling the complex thermal decomposition process is using empirical model, which uses global kinetics. The Arrhenius expression is used to correlate the rate of mass loss and temperature during the thermal decomposition.

The thermal decomposition of biomass followed the reaction as below:

\[
\text{Biomass} \xrightarrow{k} \text{Volatile} + \text{Char}
\]

The general rate of decomposition or reactivity, \(k\) can be characterised in term of temperature dependent reaction, \(k(T)\). This temperature dependent reaction obeys
the Arrhenius Eq. that is represented by activation energy \( E_a \) (J/mol) and a preexponential factor, \( A \) (s\(^{-1}\)) which expressed by:

\[
k(T) = A \exp \left( -\frac{E_a}{RT} \right)
\]

(Eq. 4.1)

where \( T \) is the absolute temperature and gas constant, \( R = 8.314\text{JK}^{-1}\text{mol}^{-1} \)

The study by Benitez (2014) described the first order reaction model as the reaction model that fit better for biomass decomposition. Therefore, this study assumed the decomposition of biomass follows the first-order reaction model. Thus, the reactivity of the decomposition over time, \( \frac{d\alpha}{dt} \) can be described in Eq. 4.2:

\[
\frac{d\alpha}{dt} = k(1 - \alpha)
\]

(Eq. 4.2)

Where the degree of conversion, \( \alpha \) or the fractional weight conversion is represented by the decomposed amount of the sample as shown in Eq. 4.3. \( w_i \) is the sample weight, \( w_o \) is the original sample weight and \( w_\infty \) is the residual or final mass after decomposition. The conversion degree, \( \alpha \) varies from 0 to 1, indicating 1 as end of conversion phase with no mass loss at 1

\[
\alpha = \frac{w_o - w_i}{w_o - w_\infty}
\]

(Eq. 4.3)

Under non-isothermal analysis, where the heating rate is maintained constant for each test, the temperature changes at a constant positive rate, \( \beta = \frac{dT}{dt} \). The rate of decomposition reaction of biomass is described as a function of temperature instead of time by combining Eq. 4.1 and Eq. 4.2 leads to Eq. 4.4 as below:

\[
\frac{d\alpha}{dT} = \frac{A(1-\alpha)}{\beta} \exp \left( -\frac{E_a}{RT} \right)
\]

(Eq. 4.4)

The Eq. 4.4 is solved using integration subject to initial condition \( \alpha = 0, \ T = T_o \) to get:

\[
\ln(1 - \alpha) = -\frac{A}{\beta} \int_{T_o}^{T} \exp \left( -\frac{E_a}{RT} \right) \, dT
\]

(Eq. 4.5)

since there is no decomposition where \( \alpha = 0 \) up to \( T_o \) (ignition temperature), the limit of the integral are conventionally changed to \( \int_{0}^{T} \exp(-\frac{E_a}{RT}) \), which leads to introduction of function \( p(x) \) as in Eq. 4.6, where \( x = \frac{E_a}{RT} \)

\[
p(x) = \int_{x}^{\infty} \frac{\exp(-x)}{x^2} \, dx
\]

(Eq. 4.6)
Thus, Eq. 4.6 reduces to:

\[
\ln(1 - \alpha) = -\frac{AE_a}{\beta R} p(x) \quad \text{(Eq. 4.7)}
\]

Based on the equation above, Doyle’s and Coats-Redfern’s approximations are adopted, due to their simplicity and easier manipulation of linear forms (Sima-Ella, Yuan, & Mays, 2005). For temperature integral approximation, Doyle suggested a linear approximation to the temperature integral, which is derived by observing a linear relationship between \(\ln p(x)\) and \(x\) as: \(p(x) \approx \exp(-5.33 - x)\). Consequently, the linear for of Eq. 4.7 can be manipulated into Eq. 4.8 as below:

\[
\ln[-\ln(1 - \alpha)] = \ln \left( -\frac{AE_a}{\beta R} \right) - 5.33 - 1.052 \left( \frac{E_a}{RT} \right) \quad \text{(Eq. 4.8)}
\]

Hence the linear Eq. can be solved for a specific heating rate, \(E\) and \(A\) can be determined from value of the slope and the intercept of plot \(\ln[-\ln(1 - \alpha)]\) versus \(1/T\). Meanwhile, in the integral approximation by Coat-Redfern, rearranging the Eq. 4.4 and integrating the temperature the Eq. 4.9 can be obtained:

\[
\ln \left( \frac{-\ln(1 - \alpha)}{T^2} \right) = \ln \left( \frac{AE_a}{\beta R} \right) - \frac{E_a}{RT} \quad \text{(Eq. 4.9)}
\]

Thus the kinetic data in this study will be interpreted using the temperature integral approximation based on the plot of \(\ln \left( \frac{-\ln(1 - \alpha)}{T^2} \right)\) versus \(\frac{1}{T}\), which will give the values of \(A\) and \(E_a\) from the intercept and slope respectively.

### 4.3.3 Kinetic parameters of the samples

The activation energy \((E_a)\) and pre-exponential factor \((A)\) for microwave torrefied sample and the non-torrefied sample of thermal decomposition for both in air and nitrogen at different heating rates are calculated based on the Doyle’s and Coats-Redfern’s approximations explained in Section 4.3.2. Relationships between the \(E_a\) and \(A\) of the biomass fuel samples can be obtained using linear regression analysis. Therefore, to show how close the data are to the fitted regression line, the coefficient of determination \((R^2)\) of the calculated linear models were also included in the presented results. The \(R^2\) calculated were between 0.9658 and 0.9989, which indicates that the linear models explain all the variability of the response data around
its mean. The equations of linear regression for kinetic parameters determination of each sample in air for is shown in Table 4.1, while Table 4.2 shows the kinetic parameters determination of each sample in nitrogen.

### Table 4.1: The equations of linear regression for kinetic parameters determination of thermal decomposition in air

<table>
<thead>
<tr>
<th>Heating rate</th>
<th>Non-torrefied sample</th>
<th>Microwave torrefied sample</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Doyle</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Coats-Redfern</td>
<td></td>
</tr>
<tr>
<td></td>
<td>10°C/min</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Linear regression</td>
<td>Linear regression</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td></td>
<td>$y = -11132x + 17.389$</td>
<td>$y = -10409x + 3.3686$</td>
</tr>
<tr>
<td></td>
<td>0.9881</td>
<td>0.9852</td>
</tr>
<tr>
<td></td>
<td>$y = -9100.4x + 13.735$</td>
<td>$y = -7763.2x - 1.0323$</td>
</tr>
<tr>
<td></td>
<td>0.9968</td>
<td>0.9959</td>
</tr>
<tr>
<td></td>
<td>$y = -9617.4x + 14.634$</td>
<td>$y = -8372.4x - 0.2348$</td>
</tr>
<tr>
<td></td>
<td>0.9975</td>
<td>0.9970</td>
</tr>
<tr>
<td></td>
<td>20°C/min</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Linear regression</td>
<td>Linear regression</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td></td>
<td>$y = -7929.2x + 12.116$</td>
<td>$y = -6719.2x - 2.6961$</td>
</tr>
<tr>
<td></td>
<td>0.9935</td>
<td>0.9915</td>
</tr>
<tr>
<td></td>
<td>$y = -8939.2x + 13.224$</td>
<td>$y = -7984.1x - 0.6896$</td>
</tr>
<tr>
<td></td>
<td>0.9945</td>
<td>0.9968</td>
</tr>
<tr>
<td></td>
<td>$y = -7603.3x + 11.005$</td>
<td>$y = -6386.1x - 3.8202$</td>
</tr>
<tr>
<td></td>
<td>0.9745</td>
<td>0.9658</td>
</tr>
<tr>
<td></td>
<td>30°C/min</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Linear regression</td>
<td>Linear regression</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td></td>
<td>$y = -8300.8x + 12.573$</td>
<td>$y = -7061.7x - 2.2874$</td>
</tr>
<tr>
<td></td>
<td>0.9990</td>
<td>0.9989</td>
</tr>
<tr>
<td></td>
<td>$y = -8764.7x + 13.087$</td>
<td>$y = -7503x - 1.8095$</td>
</tr>
<tr>
<td></td>
<td>0.9984</td>
<td>0.9980</td>
</tr>
<tr>
<td></td>
<td>$y = -8973x + 13.446$</td>
<td>$y = -7714.4x - 1.4445$</td>
</tr>
<tr>
<td></td>
<td>0.9986</td>
<td>0.9984</td>
</tr>
</tbody>
</table>

### Table 4.2: The equations of linear regression for kinetic parameters determination of thermal decomposition in nitrogen

<table>
<thead>
<tr>
<th>Heating rate</th>
<th>Non-torrefied sample</th>
<th>Microwave torrefied sample</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Doyle</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Coats-Redfern</td>
<td></td>
</tr>
<tr>
<td></td>
<td>10°C/min</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Linear regression</td>
<td>Linear regression</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td></td>
<td>$y = -8300.8x + 12.573$</td>
<td>$y = -7061.7x - 2.2874$</td>
</tr>
<tr>
<td></td>
<td>0.9990</td>
<td>0.9989</td>
</tr>
<tr>
<td></td>
<td>$y = -8764.7x + 13.087$</td>
<td>$y = -7503x - 1.8095$</td>
</tr>
<tr>
<td></td>
<td>0.9984</td>
<td>0.9980</td>
</tr>
<tr>
<td></td>
<td>$y = -8973x + 13.446$</td>
<td>$y = -7714.4x - 1.4445$</td>
</tr>
<tr>
<td></td>
<td>0.9986</td>
<td>0.9984</td>
</tr>
<tr>
<td></td>
<td>20°C/min</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Linear regression</td>
<td>Linear regression</td>
</tr>
<tr>
<td></td>
<td>$R^2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td></td>
<td>$y = -8373.3x + 12.349$</td>
<td>$y = -7123.4 - 2.528$</td>
</tr>
<tr>
<td></td>
<td>0.9942</td>
<td>0.9925</td>
</tr>
<tr>
<td></td>
<td>$y = -8469.3x + 12.323$</td>
<td>$y = -7209.7x - 2.569$</td>
</tr>
<tr>
<td></td>
<td>0.9947</td>
<td>0.9931</td>
</tr>
<tr>
<td></td>
<td>$y = -9082.3x + 12.249$</td>
<td>$y = -7804.3x - 1.677$</td>
</tr>
<tr>
<td></td>
<td>0.9970</td>
<td>0.9969</td>
</tr>
</tbody>
</table>

The activation energy ($E_a$) and pre-exponential factor ($A$) for microwave torrefied sample and the non-torrefied sample are presented in Table 4.3 and Table 4.4 respectively. Based on the results presented in Table 4.3, it can be concluded that the
kinetic parameters results determined were almost the same for both methods of calculation and in good agreement. For the non-torrefied sample, the activation energy calculated was between 64.5 to 84.4 kJ/mol, and torrefied biomass was between 53.1 to 70.7 kJ/mol. However, the results of the activation energies calculated for non-torrefied sample were found to be higher than the data reported by several researchers such as Koseki (2012) for white wood; Fisher et al. (2012) and Saddawi et al. (2010) both for raw willow.

Table 4.3: Kinetic parameters of thermal decomposition in air for samples evaluated based on Doyle’s and Coats-Redfern’s approximations

<table>
<thead>
<tr>
<th>Heating rate</th>
<th>Doyle</th>
<th>Coat-Redfern</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Eₐ (kJmol⁻¹)</td>
<td>A (s⁻¹)</td>
</tr>
<tr>
<td>10°C/min</td>
<td>84.4</td>
<td>4.40E+04</td>
</tr>
<tr>
<td>20°C/min</td>
<td>71.9</td>
<td>7.34E+03</td>
</tr>
<tr>
<td>30°C/min</td>
<td>76.0</td>
<td>2.56E+04</td>
</tr>
</tbody>
</table>

Table 4.4: Kinetic parameters of thermal decomposition in nitrogen for samples evaluated based on Doyle’s and Coats-Redfern’s approximations

<table>
<thead>
<tr>
<th>Heating rate</th>
<th>Doyle</th>
<th>Coat-Redfern</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Eₐ (kJmol⁻¹)</td>
<td>A (s⁻¹)</td>
</tr>
<tr>
<td>10°C/min</td>
<td>62.7</td>
<td>8.34E+02</td>
</tr>
<tr>
<td>20°C/min</td>
<td>70.7</td>
<td>4.48E+03</td>
</tr>
<tr>
<td>30°C/min</td>
<td>60.1</td>
<td>8.59E+02</td>
</tr>
</tbody>
</table>
Chapter 4: Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis

However, based on the results determined, the lower activation energy for torrefied material showed that the material is easier to decompose and have a higher tendency to self-heat. This result agreed with findings by Ren et al. (2013), which showed a decreasing trend of activation energy for the torrefied sample compared to the raw material. The range of kinetic parameters for thermal decomposition in nitrogen shows a higher value for both samples. For the non-torrefied sample, the activation energy calculated was between 58.7 to 70.9 kJ/mol, and torrefied biomass was between 59.2 to 71.8 kJ/mol. Therefore, it is suggested that thermal decomposition of the samples is less reactive in nitrogen.

4.4 Thermal decomposition characteristic in air

Analysis of the thermal decomposition behaviour in the air for both samples at three heating rates, 10°C/min, 20°C/min and 30°C/min are performed based on method explained in earlier in Section 4.1. Figure 4.8 shows the weight loss of non-torrefied sample with the temperature at various heating rates. As expected, we can see three regions of decomposition from the curves, where the first region from ambient temperature to 130°C that represented the moisture loss in the sample. The second region takes place from 130°C to 360°C that characterised by volatilization process of the cellulosic and hemicellulose components. Lastly, the third region from varies between 360°C to 900°C is characterised by the char combustion, which produced the volatiles and solid residues.

![Figure 4.8: Temperature profile of decomposition in air of non-torrefied sample at different heating rates](image-url)
The clear stages are due to the different compositions of biomass that degraded at a different temperature. The non-torrefied sample is originated from the biomass fuel that did not undergo thermal pre-treatment. Therefore all the compositions of the biomass are still within the sample. The composition of volatile matters also higher in the non-torrefied sample. The higher volatile matters cause the non-torrefied sample to burn at a lower temperature than microwave torrefied sample.

Based on Figure 4.9, the noticeable second peak for a non-torrefied sample during the decomposition process can be seen. The first peak is associated with the main devolatilisation stage and the second peak represents the char oxidation. The non-torrefied material showed a significant second peak, which suggested that there is volatiles component in the material such as hemicellulose, cellulose and lignin that decomposed at a different temperature.

Figure 4.9: Temperature profile of derivative weight loss in air of non-torrefied sample at different heating rate

Figure 4.10 show that the thermal decomposition in of microwave torrefied sample does not have a clear first peak of moisture loss. There is no clear peak detected due to the small amount of moisture content from the sample. The torrefaction process managed to reduce the water content greatly and retained the energy in solid mass. The same pattern of temperature profile can be found for each heating rate, where a
clear devolatilization stage can be witnessed at the second phase of the heating profile, which is between 280 °C – 450 °C.

![Graph showing weight loss vs. temperature](image1.png)

**Figure 4.10:** Temperature profile of decomposition in air for microwave torrefied sample at different heating rate

There is no clear indication for the second peak in the case of the microwave torrefied sample as shown in Figure 4.11. This pattern is found in the temperature profile of the microwaved torrefied sample due to the moisture loss during the torrefaction process. Besides that, it can be seen that the maximum weight loss temperature of the sample shifted to higher temperature when the heating rate increase from 10 °C/min to 30 °C/min.

![Graph showing derivative weight loss vs. temperature](image2.png)

**Figure 4.11:** Temperature profile of derivative weight loss in air for microwave torrefied sample at different heating rate
Based on Figure 4.8 and Figure 4.10, it can be observed that with the increase of heating rate, the curves move slightly to a higher temperature zone, for both samples. The delay of the temperature due to several factors such as shorter reaction time and lower degree of reaction changes due to higher heating rate (Guo et al., 2012), thus the temperature required for the sample to reach the same conversion will be higher (Heydari et al., 2015). Besides that, the low thermal conductivity of the samples can also contribute to this phenomenon where the longer time needed for the heat to transfer across the sample and there is no enough time available for the heat to transfer within the sample matrix (Kan, Strezov, & Evans, 2016).

Figure 4.10, also shows that microwave torrefied sample generated a larger amount of unburned residual material at the end of the experiment compared to the non-torrefied sample in Figure 4.8. This is in accordance with the higher ash content in microwave torrefied sample. However, non-torrefied sample decomposed in at 30°C/min generated more residual material compared to the one at 10°C and 20°C, which could be due to the shorter reaction time with the low degree reaction changes as mentioned earlier.

### 4.5 Thermal decomposition characteristic in nitrogen

Thermal results obtained from decomposition in nitrogen for the non-torrefied sample are presented in Figure 4.12 and Figure 4.13. Based on the decomposition in nitrogen for both samples, the residual matter at the end of the reaction is higher than the one done in air. The higher amount of residual matter is expected due to the incomplete decomposition in nitrogen over air. The Figure 4.12 shows the temperature profile of the non-torrefied sample heated at various heating rate. It can be witnessed that, different heating rates did not have strong influence on the amount of residual left at the end of the heating process of the samples in nitrogen. For non-torrefied sample, the residual left at the end of process is 16% of total weight of sample.
Figure 4.12: Temperature profile of decomposition in nitrogen of non-torrefied sample at different heating rate

It was examined that the decomposition proceeds through three stages of the sample mass loss. The first stage is assigned to the moisture loss through the water evaporation, while the second stage, the mass loss is due thermal decomposition of wood compounds, that consist of hemicellulose and cellulose. The last stage is the decomposition of lignin that decomposed in the wider temperature range up to 900°C.

The same pattern of heating rate effect can be witnessed during decomposition in nitrogen for non-torrefied sample in Figure 4.13. It can be witnessed that, the reaction time reduces and the temperature profiles of the reaction move to the higher temperature zone. The same behaviour was found by other researchers such as Cruz Ceballos et al. (2015); Benitez (2014); Guo et al. (2012); Gašparovič et al. (2010). However, compared to the thermal decomposition in air, there is no third peak can be observed for both samples when decomposed in nitrogen.
Chapter 4: Experimental determination of reactivity and kinetic parameters using thermogravimetric analysis

Figure 4.13: Temperature profile of derivative weight loss in nitrogen non-torrefied sample at different heating rate

The same behaviour can also be found in the decomposition of microwave torrefied sample in nitrogen as shown in Figure 4.14. The unburned residual for microwave torrefied sample is higher compared to the non-torrefied sample, due to the changes of properties of the microwave torrefied sample that having lower volatiles and higher ash content. Therefore, more residual is observed due to pyrolytic decomposition of the sample. The amount of the residual left at the end of the heating process was observed at 25% of the total weight of sample.

Figure 4.14: Temperature profile of decomposition in nitrogen of microwave torrefied sample at different heating rate
The same pattern of heating behaviour as non-torrefied sample can be seen during the analysis of the microwave torrefied sample in nitrogen at various heating rate as shown in Figure 4.15. There is a clear second peaks for the samples heated at various heating rates. However the peaks tend to shift to a higher temperature, with the decrease of heating rates.

![Figure 4.15: Temperature profile of derivative weight loss in nitrogen for microwave torrefied sample at different heating rate](image)

**4.5.1 Characteristic parameters of both samples at various heating rate and carrier gas**

Table 4.5 presents the comparison of characteristic parameters and maximum rate of mass loss of the decomposition in air and nitrogen. It is not much difference between the temperatures of initial combustion (TIC) for thermal decomposition of the non-torrefied sample in air and nitrogen. However, the trend of characteristic temperatures increase can be seen for microwave torrefied sample comparing to the nitrogen. The increase in the temperature when decomposed in nitrogen is due to the reduction in decomposition rate. The maximum weight losses of both samples heated in air are much lower than the one heated in nitrogen due to its pyrolytic condition.
Table 4.5: Characteristic parameters and maximum rate of mass loss (DTG$_{max}$) for the decomposition in air and nitrogen

<table>
<thead>
<tr>
<th>Sample</th>
<th>Heating rate</th>
<th>TIC (°C)</th>
<th>TMWL (°C)</th>
<th>DTG$_{max}$ (%/min)</th>
<th>TIC (°C)</th>
<th>TMWL (°C)</th>
<th>DTG$_{max}$ (%/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-torrefied sample</td>
<td>10°C/min</td>
<td>305</td>
<td>350</td>
<td>15.19</td>
<td>305</td>
<td>378</td>
<td>11.35</td>
</tr>
<tr>
<td></td>
<td>20°C/min</td>
<td>310</td>
<td>360</td>
<td>19.17</td>
<td>310</td>
<td>387</td>
<td>17.32</td>
</tr>
<tr>
<td></td>
<td>30°C/min</td>
<td>315</td>
<td>374</td>
<td>23.75</td>
<td>320</td>
<td>386</td>
<td>25.49</td>
</tr>
<tr>
<td>Microwave torrefied sample</td>
<td>10°C/min</td>
<td>305</td>
<td>346</td>
<td>8.03</td>
<td>325</td>
<td>376</td>
<td>8.30</td>
</tr>
<tr>
<td></td>
<td>20°C/min</td>
<td>325</td>
<td>369</td>
<td>16.43</td>
<td>330</td>
<td>384</td>
<td>15.78</td>
</tr>
<tr>
<td></td>
<td>30°C/min</td>
<td>330</td>
<td>374</td>
<td>23.79</td>
<td>335</td>
<td>385</td>
<td>22.97</td>
</tr>
</tbody>
</table>

4.6 Evaluation of self-ignition risk based on characteristic oxidation temperature and activation energy

Ramírez et al., (2010) have described the degree of susceptibility of material that prone to self-ignition can be evaluated based on the relationship of the characteristic oxidation temperature (T$_{char}$) and the activation energy (E$_a$). The risk assessment as the function of characteristic oxidation temperature and activation energy for coals have been explained in Krause (2009). Where the risk has been classified into four groups, which are: (i) very high risk, (ii) high risk, (iii) medium risk and lastly (iv) low risk.

For the comparison purposes, the value of the parameters determined this study have been plotted among the data from García Torrent et al. (2016) and Ramírez et al. (2010) with the proposed classification groups from Krause (2009). Based on Figure 4.16, risk ranking graph plotted the present study data is written in red, and the data from other researchers is written in black. Based on the plotted data, the non-torrefied sample and microwave torrefied sample are classified as material with medium risk. They fall in the same group as other lignocellulosic biomass such as straw, coconut, olive residue and pine cone.

From the plotted graph, it is clear that the location of the microwave torrefied sample fall in the lower region of the plotted data, which indicates that it has a higher tendency to self-ignite compared to the non-torrefied sample due to its lower
activation energy. Besides that, it can be witnessed that the samples tested in this study had a higher risk compared to bituminous coal and sub-bituminous coal. In addition to that, biomasses such as wood chip and animal waste are easier to self-ignite compared to the samples used in this study.

Figure 4.16: Risk ranking graph based on the activation energy and characteristic oxidation temperature plotted among data from García Torrent et al., (2016) and Ramírez et al., (2010)

4.7 Summary

- The thermogravimetric curves move towards higher temperature region when torrefied biomass sample is decomposed as it suggested that the material is more stable towards thermal decomposition.
- The result of TIC and TMWL for both samples showed that the non-torrefied sample is more reactive than the torrefied sample. However, $T_{\text{charac}}$ for the non-torrefied sample is higher than the torrefied sample.
- Thermal decomposition in nitrogen for both samples shows higher residual matter at the end of the reaction is compared to the one done in air. This is expected due to the incomplete decomposition in nitrogen over air.
- The activation energy for thermal decomposition in air is in between 64.5 to 84.4 kJ/mol for the non-torrefied sample, and 53.1 to 70.7 kJ/mol torrefied.
sample. While in nitrogen is in between 58.7 to 70.9 kJ/mol for non-torrefied sample and 59.2 to 71.8 kJ/mol for torrefied sample.

- The lower activation energy for torrefied material showed that the material is easier to decompose, as a result, have a higher tendency to self-heat.
- The kinetic parameters findings agreed to the results found in the $T_{\text{charac}}$ analysis, which confirmed that the microwave torrefied sample is more reactive than non-torrefied sample in air.
- Based on the risk ranking graph; both non-torrefied sample and microwave torrefied sample are classified as a material with medium risk. The location of plotted point for microwave torrefied sample fall in lower part of the graph, which indicates that it has a higher tendency to self-ignite compared to the non-torrefied sample due to its lower activation energy.
Chapter 5 Experimental study of the thermal runaway in bulk tests

Chapter 5 provides details of experiments undertaken the heating behaviour of the samples heated up to various bulk sizes and oven temperatures. The experimental programme considers examining the effect of the bulk size and the oven temperature on time taken for a thermal runaway to occur. This chapter presents the temperature profile from each experiment. This chapter also shows the relationship between the ignition delay time and oven temperature as well as the bulk size thermal runaway to happen in a high-temperature environment.

5.1 Experimental works

5.1.1 Objectives of the experiment
The experimental works done in this chapter is to fulfil the objectives as below:
(i) to determine the relationship between various bulk sizes on the ignition delay time for the ignition to occur;
(ii) to determine the relationship between the various oven temperatures and the time taken for a thermal runaway to occur; and
(iii) to compare the thermal stability of both materials in larger scale heating environment

5.1.2 Samples and equipment
The samples used in this experiment are non-torrefied sample and microwave torrefied sample. The properties of both samples are very different from one another. Therefore, it will give a good comparison of heating behaviour. The experiments were conducted using the various size of cubic shaped wire mesh bulks as shown in Figure 5.1 with each cube's length labelled and named as Bulk 1, Bulk 2 and Bulk 3. The schematic diagrams of the experimental setup with the position of the thermocouples are presented in Figure 5.2, Figure 5.3 and Figure 5.4.
Figure 5.1: Three sizes of wire mesh baskets wide side lengths labelled

Figure 5.2: Schematic diagram of experimental setup of heating tests for Bulk 1
Figure 5.3: Schematic diagram of experimental setup of heating tests for Bulk 2

Figure 5.4: Schematic diagram of experimental setup of heating tests for Bulk 3
5.1.3 Thermal data logging system

The temperature measurements were made using five type-K thermocouples with Inconel sheath (±2.5°C tolerances according to BS 4937) The data from the thermocouple we collected using Microlink 751 data logging system (Microlink Engineering Solution, Manchester, England). The data logger was connected to a laptop and recorded temperature every second. The data logger system continuously recorded the temperatures, while the data acquisition software (Windmill software) stored the input data. The data were transferred to Microsoft Excel for further analysis. The thermocouple readings for each bulk test were presented in the Appendix A to display the raw temperature measurements captured by each thermocouple during the test.

5.2 Experimental procedure

This experimental works focused on self-heating phenomena of two types of bulk samples that lead to thermal runaway. The samples used in these experiments are non-torrefied sample and the microwave torrefied sample as discussed in Chapter 3. The experiments were carried out to investigate the heating behaviour of the sample using three different bulk sizes subjected to several constant oven temperatures.

The experiments were performed using a laboratory oven (Carbolite, LHT 6/30). The oven has a capacity of 30 L with an internal dimension of 300 x 300 x 305 mm (length x width x depth). The oven is thermostatically controlled and equipped with an air circulation fan that helps to reduce the temperature difference inside the oven chamber.

The open-top wire mesh basket acted as a reactor. The wire mesh basket was made from stainless steel of 0.25μm diameter mesh. The wire mesh basket was placed inside the oven and thermocouples were properly positioned according to the coordinate shown in Table 5.1. The experiments were repeated for three different sample volumes, Bulk 1 (216cm$^3$), Bulk 2 (512 cm$^3$) and Bulk 3 (1000 cm$^3$), with dimensions stated in Table 5.1.
Chapter 5: Experimental study of the thermal runaway in bulk tests

Figure 5.5 shows the positions for the thermocouples inside the bulk, which are T1 in the middle, T2 halfway from the centre to the wall (radial), T3 at the wall of the bulk and T4 at the opposite wall of the bulk. Lastly, T5 is for oven temperature measurement. The tips of thermocouples were kept at the same axial level from the bottom. The locations of the tips for each sample volume are based on their vertical and radial coordinates as shown in Table 5.1. The temperatures were being recorded by a data acquisition system.

Table 5.1: Locations of the thermocouples based on vertical and radial coordinates

<table>
<thead>
<tr>
<th></th>
<th>Length, L (mm)</th>
<th>Height, H (mm)</th>
<th>Width, W (mm)</th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>T4</th>
<th>T5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk 1</td>
<td>60</td>
<td>60</td>
<td>60</td>
<td>(0,0)</td>
<td>(30,15)</td>
<td>(30,30)</td>
<td>(30,-30)</td>
<td>Oven</td>
</tr>
<tr>
<td>Bulk 2</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>(0,0)</td>
<td>(40,20)</td>
<td>(40,40)</td>
<td>(40,-40)</td>
<td>Oven</td>
</tr>
<tr>
<td>Bulk 3</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>(0,0)</td>
<td>(50,25)</td>
<td>(50,50)</td>
<td>(50,-50)</td>
<td>Oven</td>
</tr>
</tbody>
</table>
After the thermocouples had been properly positioned and the wire mesh basket was suspended at the centre of the oven, the samples were fully filled into the wire mesh basket. Figure 5.6 shows the example of wire mesh basket that is fully filled with microwave torrefied sample. The oven temperature was maintained at a selected constant temperature.

Figure 5.6: Wire mesh basket filled with microwave torrefied sample

The samples were subjected to the selected temperature until the reading of the thermocouple at the centre (T1) exceeded the oven temperature (T5). The exceeded temperature is an indication that self-heating had occurred in bulk followed by the ignition. If the ignition occurred, the oven is turned off, and the samples were let to burn fully. The oven was then cooled down to room temperature before the next batch is set up using the same method. The full list of a bulk heating experiment done is presented in Table 5.2.

Table 5.2: List of oven temperatures used during bulk heating experiments

<table>
<thead>
<tr>
<th>Sample</th>
<th>Bulk 1</th>
<th>Bulk 2</th>
<th>Bulk 3</th>
<th>Bulk 1</th>
<th>Bulk 2</th>
<th>Bulk 3</th>
<th>Bulk 1</th>
<th>Bulk 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-torrefied</td>
<td>180°C</td>
<td>200°C</td>
<td>200°C</td>
<td>200°C</td>
<td>200°C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Torrefied</td>
<td>180°C</td>
<td>180°C</td>
<td>180°C</td>
<td>170°C</td>
<td>190°C</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
5.3 Self-heating and thermal runaway in bulk tests

The bulk experiment test involved boundary heating of an initially cold exothermic material exposed to a hot environment with a constant temperature. This experiment is adapted from basket heating method described by Chen (1999), which intended to determine the kinetic analysis of self-heating process. However, this experimental work is anticipated to examine the self-heating propensity of the samples without the consideration of its kinetics. The heating behaviours of the sample were investigated using various cubic shape wire mesh bulk, as shown in Figure 5.1. The bulk was filled with the samples, and the constant oven temperature was maintained throughout the experiment.

The experimental works are designed to investigate the thermal stability of the samples based on the bulk size and the oven temperature, by comparing the induction time of the sample to ignite. The induction time in this experimental works is defined as the time taken for the centre temperature (T1) to exceed the oven temperature (T5). Based on the discussion in Section 2.5.2, the temperature profile of the samples followed the supercritical curves when self-ignition occurred in the system. The supercritical state represents that the sample reaches the critical temperature, which the self-heating follows by the thermal runaway occur. Meanwhile, if there is no self-heating detected, the temperature profile follows the subcritical curve. The subcritical curve indicates that the heat generation is equal to the heat dissipation. Thus equilibrium state with no rapid increases of temperature is achieved.

5.3.1 Discussion of the self-heating propensity of the samples

Based on the kinetic study of the biomass fuels presented in Section 4.3, the non-torrefied sample has higher apparent activation energy compare to the microwave torrefied sample. By doing this set of experiment, the heating behaviour of the fuel in larger scale can be examined. Thus, in order to confirm the results, the bulk test is performed at an oven temperature of 180°C using wire mesh Bulk 1 (side length 60mm) for both samples.
Figure 5.7 shows the temperature profiles of T1 (thermocouple at the centre) for both samples. The rapid temperature rise can be seen in the temperature profile of microwave torrefied sample. This rapid temperature rise exhibits that the microwave torrefied sample experienced self-heating stage that leads to the thermal runaway. The temperature profile follows the supercritical behaviour, where the heat production exceeded the heat loss of the system that result in unsteady state. In contrast to the microwave torrefied sample, the curve for non-torrefied sample shows the subcritical behaviour where the sample become hotter approaching oven temperature but no ignition observed.

The observation from the heating tests confirmed the findings of the kinetic parameter of the samples, where at the same heating condition, self-heating that lead to thermal runaway can only be detected for microwave torrefied sample. Whereas, there is no self-heating phenomenon detected for the non-torrefied sample. This observation confirms that the microwave torrefied sample is more reactive compared to the non-torrefied sample.

![Temperature Profiles](image)

**Figure 5.7: The comparison of the temperature profiles of the samples at 180°C heated using Bulk 1**

We will move on to discuss further temperature profiles obtained for each sample. There are significant different between the temperature profile during heating process of the non-torrefied sample and the microwave torrefied sample.
5.3.1.1 Non-torrefied sample

The temperatures profile reached it plateau after 6700 seconds (after black dotted line). It can be witnessed that after 6700 seconds the thermocouple's reading at the centre of the sample, T1 follows the same temperature pattern as oven temperature, T5. There are no self-heating detected after more than 4 hours (15000 seconds), which indicated that the system achieved steady state and subcritical behaviour is observed. The temperature of the whole sample remains constant throughout the experiment when it reached the oven temperature. The measured temperature profiles of the event are shown in Figure 5.8.

![Graph showing temperature profiles](image)

**Figure 5.8: Temperature profile of non-torrefied sample heated at 180°C in Bulk 1**

In other words, the surrounding temperature balances the temperature generated by the exothermic heat generation of the non-torrefied sample in bulk. Therefore no thermal runaway occurred. Figure 5.9 gives clearer view of the temperature profile of T1 and T5 (oven). The temperature of T exceeded oven temperature, but with only small temperature differences of 3°C, with the highest recorded temperature at T1 was 183°C. Thus, there was minimal heat generated by the system, and the temperature rise was able to be balanced by the surrounding temperature. Therefore, no self-heating detected, due to the size of bulk used, which is 60 mm side length. The short side length, allowed the heat to disperse to the surrounding easily and balanced the heat transfer in the system.
Figure 5.9: Temperature profile of T1 and oven temperature (T5)

5.3.1.2 Microwave torrefied sample

The same heating condition applied to the microwave torrefied sample (oven temperature 180°C heated in Bulk 1. The comparison of the temperature profiles for T1, T3 and T5 thermocouples are shown in Figure 5.10, while the temperature reading for all the thermocouples shown in Appendix A-1. Following the experimental technique, the comparisons of those thermocouples reading are significant in the observation of the self-heating behaviour.

Figure 5.10: Temperature profile of microwave torrefied sample at 180°C in Bulk 1
Based on Figure 5.10, the temperature of the centre (T1) becomes higher than the temperature at the edge of bulk (T3) and oven temperature (T5) indicates that the self-heating phenomena followed by thermal runaway occurred in the system. The microwave torrefied sample started to self-heat at 4002 seconds (66.7 minutes), where the T1 reading indicating rapid increases in temperature and thermal runaway is detected at this stage. The rapid growth is the indicator that the heat generated by the exothermic reaction of the material is more than the heat dissipated to the surrounding. At this point, it can be established that a self-heating process had started and followed by the self-ignition of the sample.

Consequently, by comparing the behaviour of both samples, it can be established that microwave torrefied sample is more reactive towards heat compared to non-torrefied sample. This finding also in agreement to the kinetic analysis results discussed in Section 4.3, which the apparent activation energy of the microwave torrefied sample is lower than the non-torrefied sample. Therefore, the heating behaviour reflected that the thermal stability of microwave torrefied sample is less than the non-torrefied sample.

5.3.2 Discussion of effect of bulk size on self-heating behaviour

a) Non-torrefied sample

The effect of different bulk sizes is tested for non-torrefied samples, where the samples were heated in the cubic shape bulk with different volume Bulk 1 (216cm³), Bulk 2 (512cm³) and Bulk 3 (1000cm³) and the oven temperature was set at 200°C for each experiment. Based on the experiments, it can be seen that the temperature profile of T1 for the samples shifted to the longer induction time as the volume is larger as illustrated in Figure 5.11. The temperature reading of T1 for non-torrefied sample heated in a largest cubic volume of 1000 cm³, showed that the longest time taken before it reached its critical stage. While T1 for the non-torrefied sample heated at the same oven temperature in smallest cube taken shortest to reached its critical stage.
Figure 5.11: Comparison of temperature reading at the centre (T1) of non-torrefied sample heated at 200°C at different volumes

The shift of temperature profile is due to the longer time taken of the heat from the surrounding to dissipate to the centre of the bulk, as the size is larger. The heating process of the whole bulk is taking longer for greater volume. The pattern follows the pattern of previous researchers that shows the critical storage time correlate linearly with the storage volumes in their work (García-Torrent et al., 2012; Ramírez, García-torrent, & Tascón, 2010; Yan et al., 2005).

The experiment managed to establish the relationship between the bulk size and the induction time. The data obtained in Figure 5.11 can be found from the temperature profile of each test. The induction time for each bulk size is 5767 seconds for Bulk 1 (as shown in Figure 5.12), 8159 seconds for Bulk 2 (as illustrated in Figure 5.13), and 11487 seconds for Bulk 3 (as illustrated in Figure 5.14). The complete thermocouple readings of the tests can be seen in Appendix A-2 for Bulk 1, Appendix A-3 for Bulk 2 and lastly Appendix A- 4 for Bulk 3.
Chapter 5: Experimental study of the thermal runaway in bulk tests

Figure 5.12: Induction time of non-torrefied sample heated at 200°C in Bulk 1

Figure 5.13: Induction time of non-torrefied sample heated at 200°C in Bulk 2

Figure 5.14: Induction time of non-torrefied sample heated at 200°C in Bulk 3
A linear graph had been plotted using the information of induction time and volume of the bulk used for heating test. The linear graph to show the relationship between the volume and the induction time of the non-torrefied sample heated at 200°C is shown in Figure 5.15. It can be concluded that, the larger the volume; the longer time taken for the samples to reach its critical stage. Much longer time to achieve supercritical condition indicating that it takes longer time for the heat to transfer to the centre of the bulk to trigger the ignition.

![Figure 5.15: Induction time vs V/A of non-torrefied sample at 200°C](image)

The linear relationship of the side length and the induction time is shown in Figure 5.16. It follows the same pattern linear relationship as the relationship between induction time and V/A, as the side length represent the size of the bulk.

![Figure 5.16: Relationship between side length and induction time of non-torrefied sample heated at 200°C](image)
b) Microwave torrefied sample

To examine the effect of bulk size on the microwave torrefied sample, the sample had been heated at 180°C in various bulk volumes. For this tests, lower oven temperature had been chosen due to the fact that microwave torrefied sample is more reactive compared to the non-torrefied sample. Figure 5.17 shows the induction time for microwave torrefied sample heated up in Bulk 1. The temperature profile of T1 started to heat up and slowly reaching towards the oven temperature at 180°C. Then at 4002 seconds, the centre of the sample started to self-heat and followed by a rapid temperature increased afterwards. The rapid temperature increased showed that the sample had undergone a thermal runaway.

![Figure 5.17: Temperature profile of microwave torrefied sample heated at 180°C Bulk 1](image)

Figure 5.18 shows the induction time for microwave torrefied sample heated up in Bulk 2. The temperature profile of T1 started to heat up and slowly reaching towards the oven temperature at 180°C. However, the time taken for the sample in Bulk 2 to self-heat was longer that Bulk 1. The induction time for the Bulk 2 was 5066 seconds. The self-heat also followed by a rapid temperature increased afterwards. The rapid temperature increased showed that the sample had undergone a thermal runaway.
Figure 5.18: Temperature profile of microwave torrefied heated at 180°C in Bulk 2

Lastly, Figure 5.19 shows the induction time for microwave torrefied sample heated up in Bulk 3. For this test, the induction time was the longest compared to the setup explained earlier for Bulk 1 and Bulk 2. The induction time for the Bulk 3 was recorded at 7611 seconds. A thermal runaway also detected for this test.

Figure 5.19: Temperature profile of microwave torrefied sample heated at 180°C in Bulk 3
The thermocouples readings for each mentioned tests earlier can be found in The complete thermocouple readings of the tests can be seen in Appendix A-5 for Bulk 1, Appendix A-6 for Bulk 2 and lastly Appendix A-7 for Bulk 3. Based on the above temperature profile, it can be seen that the temperature profile of T1 for the samples shifted to the longer induction time. For easier comparison, Figure 5.20 compared the temperature readings of T1 for each bulk size. The temperature reading of T1 for microwave sample heated in a largest cubic volume of 1000 cm$^3$, showed that the longest time taken before it reached its critical stage. While T1 for the microwave torrefied sample heated at the same oven temperature in smallest cube taken shortest to reached its critical stage. The same pattern can also be observed in non-torrefied sample tests.

![Figure 5.20: Comparison of temperature profiles at the centre of the bulk (T1) of microwave torrefied sample heated at 180°C at different volumes](image)

Therefore, based on the time induction for each bulk size, a linear graph was plotted. The same relationship between the size of the bulk and induction time can be observed. Figure 5.21 shows the relationship between the side length of the bulk and the induction time. The side lengths in the graph represent the size of the bulk. Therefore a liner relationship between the size of bulk and the induction time was found to follow the same pattern as the non-torrefied sample. The finding also in
agreement with the studies done by Graham (2015); Saddawi et al. (2013); Pauner & Bygbjerg (2005) where the larger the bulk size the longer the induction time.

![Graph showing relationship between side length and induction time for microwave torrefied sample heated at 180°C]

**Figure 5.21:** Relationship between side length and induction time for microwave torrefied sample heated at 180°C

### 5.3.3 Discussion of effect of oven temperature on self-heating behaviour

Various oven temperature had been chosen to examine the effect of ambient temperature towards the tendency of self-heating of the samples. The ambient temperature played an important role towards the induction time of the sample before self-heating was detected during the experiments. Thus, to examine the behaviour of the microwave torrefied sample in various oven temperatures, smallest size of the bulk was chosen. The smaller size of bulk will resulted in a higher ambient temperature need for the sample to self-heat (Veznikova et al., 2014; Wolters et al., 2003).

The samples were heated at oven temperature of 170°C, 180°C and 190°C with the same size of the cubic bulk of 216 cm³. Figure 5.22 shows the temperature profile of the microwave torrefied sample heated at 170°C. Self-heating was detected at 4440 seconds where the reading of thermocouple T1 started to exceeded the oven temperature.
However, it can be observed in Figure 5.22 that the reaction did not reach a critical state, where the temperature reading of T1 plateaued at 200°C. While Figure 5.23 shows the temperature profile of the microwave torrefied sample heated at 190°C. Conversely, for this oven temperature, self-heating was detected at earlier compared to oven temperature of 170°C, which is at 3513 seconds. The same pattern was observed during the self-heating test of refuse paper and plastic fuel by Koseki (2012), which showed that the shorter time taken for the sample to achieve the supercritical stage which indicated self-heating occurred followed by thermal runaway.

---

**Figure 5.22:** Temperature profile of microwave torrefied sample at 170°C in Bulk 1

**Figure 5.23:** Temperature profile for microwave torrefied sample at 190°C in Bulk 1
The comparison of the thermocouple readings of T1 is shown in Figure 5.24. The temperature profiles of the T1 shifted to the lower region of the graph as the higher oven temperature applied. It can be observed that the curve of sample heated at 190°C showed a substantial supercritical pattern. Whereas, for sample heated at 170°C, the curve is smoother and follows the pattern of critical heating behaviour. Based on the graph, it can be witnessed that a supercritical stage was achieved when the sample heated at 190°C and a steady state was achieved when sample heated at lower oven temperature of 170°C. The same pattern was found in the study of self-heating of coal dust accumulations by Wu & Bulck (2014), where the steady state can be witnessed for the lower oven temperature and shifted to critical state at the higher oven temperature.

Figure 5.24: Comparison of readings at the centre of the bulk (T1) of microwave torrefied sample heated in Bulk 1 at various oven temperatures

Using the information collected from the heating test, a linear graph had been plotted to show the relationship between the oven temperature and the induction time. Figure 5.25 shows an inverse relationship between the oven temperature and the induction time is established. A shorter induction time is observed for the sample heated at a higher oven temperature. The shorter reaction is predictable due to the chemical reaction that based on Arrhenius equation where the kinetic constant increases as the temperature increases. Therefore, when the sample heated at higher temperature, shorter induction time needed for the sample to self-heat.
5.4 Conclusion

- The oven temperature used for non-torrefied sample is greater than 180°C as the lower temperature does not give a sufficient heating that lead to self-heat. While at 180°C, the microwave torrefied sample showed self-heating phenomena that led to thermal runaway.
- The microwave torrefied sample is more reactive compared to the non-torrefied sample. Thus, agreeable to the findings of apparent kinetic parameters.
- The induction time increased with the increased of the bulk volume. While the induction time decreased when the oven temperature increased.
- The inversed relationship between oven temperature and induction time was found from the test.
- The same pattern of the relationship between the induction time and the bulk volume as well as the ignition time and the oven temperature is observed when non-torrefied sample is used.
Chapter 6 Development of numerical models to simulate the self-heating process in bulk storage area

This chapter details the work undertaken in the development of self-heating model based on mass and energy transfer equations for bulk storage using the properties and the kinetic parameters determined from the experiments. The comparisons of the thermal behaviour of both samples are based on the temperature profiles predicted during the self-heating process. The model was used to predict the critical ambient temperature and induction time of the self-heating process leading to ignition. The effect of height of bulk and the ambient temperature of the storage as well as properties of the material were investigated. Lastly, the model was verified by comparing the temperature profiles of the bulk scale experiments to the simulations results.

6.1 Self-heating of biomass fuels in open storage area

Self-heating within the biomass material causes the temperature rise during its storage phase. The self-heating process leading to ignition is a problem in the biomass fuels handling and storage. Self-ignition process of bulk biomass is started when the heat produced inside the pile through exothermic process unable to be dissipated to the surroundings. This situation is highly concerned when it comes to the storage of large amount of biomass that can lead to uncontrolled positive heat feedback that can lead thermal runaway.

Figure 6.1: Heat loop that leading to thermal runaway
Additional heat source may be precursor to the self-heating, such as physical processes and biological activities. However, these models only consider the exothermic reaction of chemical process. The biomass fuel is assumed to have a very low moisture content that inhibit the growth of microorganisms, therefore the exothermic reaction due to biological activities is ignored. In addition, heat from the water adsorption/condensation is not included in the model as the reaction happens at low temperature.

This work considers the heating process in rectangular shape pile, where the height is the critical parameters for simulations. The pile is assumed to be stored indoor without any aeration thus forced convection is negligible. The model of self-heating in biomass pile is solved using COMSOL Multiphysics® software Ver. 4.4. For the purpose of validation, the simulation of the bulk test is done to compare with the simulations results of this section. The parameters to be studied are the height and ambient temperature. This is because both are the easier parameter to be controlled by the workers during handling and storage phase. Thus, it is practical to study these parameters to give a clearer indications of what can the workers can do to ensure safe handling of the biomass fuels.

6.2 The model set-up

To simplify the model, a two-dimensional model of heat and mass transfer is used for the simulation. The bulk pile is deposited on a concrete base plate that was considered as the insulated wall where the heat conduction between the bulk pile and storage floor was ignored. Figure 6.2 shows the two-dimensional bulk geometry model used for the simulation.

The boundary conditions are set as the constant temperature at the beginning of the process that represents the critical ambient temperature \( T_o \) of the storage condition. The top surface of the bulk geometry is subjected to a convection boundary condition between the hotter ambient conditions that and the lower temperature of the bulk pile. The natural convection is the dominant mechanism of oxygen access into the bulk pile. The top and side surfaces were considered as open boundaries. The heat

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source \( (Q_{\text{source}}) \) term is defined using the Arrhenius type rate equations. The kinetic parameters are measured during the experimental work presented in Section 4.3.

![Figure 6.2: Geometry of simplified model of the pile in an open storage.](image)

### 6.3 Governing equations and computational module

A two-dimensional model is developed to simulate the self-heating process of both samples examined earlier in the open storage environment. The mathematical model based on the physic modules in COMSOL Multiphysics® to simulate the process of heating up and the potential of self ignition processes in the open storage stockpile. The stockpile is considered as a reactive porous medium where the prediction of possible self-ignition risk was performed by modelling the equations for heat and mass transfer, which includes the heat source from the exothermic chemical reaction.

In COMSOL Multiphysics® the numerical model of the self-heating process in the bulk piles is using a multi-physics coupled heat and mass transfer in porous media. In self-heating of biomass, the dominant source of heat is the chemical oxidation, thus based on Blomqvist & Persson (2003) it can be governed fundamentally by the rates of diffusion and convection of air from the surrounding area.
6.3.1 Mathematical model

Considering the complexity nature of the self-heating process, several assumptions had been adopted to the model in order to simplify the simulation. Thus, the following assumptions were chosen for this work:

i. The model proposed sufficiently porous and isotropic as well as a homogenous pile;

ii. The deformation, shrinkage or collapsing of the pile are not considered;

iii. The ambient temperature is considered constant throughout the simulation;

iv. Heat transfer through the body is by conduction in porous media with air filling the pores;

v. Heat transfers at the surface to the surroundings are by convection and radiation;

vi. The storage floor is a perfectly concrete slab, which perhaps leads to maximum heat build-up within the stored material;

vii. Natural convection is considered for the simulation as the stockpile storage area is assumed not to be equipped with any fan or aeration equipment;

viii. The stockpile is reactive so that the solid percentage volume and gas percentage volume remain unchanged during the self-heating; and

ix. The effect of moisture in the pile is neglected as the moisture content of the material studied is considered low and the temperature increase due to moisture is significantly low.

The governing equation of self-heating is based on the theory of energy conservation in heat transfer in the porous pile as shown in Eq. 6.1 on \(0 \leq y \leq h\):

\[
\left[ (\varepsilon \rho_g C_g) + (1 - \varepsilon) \rho_s C_s \right] \frac{\partial T}{\partial t} + \rho_g C_g \left( \frac{\partial T}{\partial y} \right) = \lambda_{eff} \left( \frac{\partial^2 T}{\partial y^2} \right) + (1 - \varepsilon) A O_2 \exp \left( - \frac{E_a}{RT} \right)
\]

(Eq.6.1)

The effective thermal conductivity in the porous media;

\[
\lambda_{eff} = \varepsilon \lambda_g + (1 - \varepsilon) \lambda_s
\]

(Eq. 6.2)

In the case of porous media, the oxygen diffusivity is considered as; \(D_{o,eff} = \varepsilon D_{o,g}\)
Thus the transport of oxygen concentration on $0 \leq y \leq h$

$$
\varepsilon \frac{\partial O_2}{\partial t} = D_{o,eff} \left( \frac{\partial^2 O_2}{\partial y^2} \right) - (1 - \varepsilon)O_2 A\exp \left( -\frac{E_a}{RT} \right)
$$

(Eq. 6.3)

An Arrhenius equation is used to represent the dependence of the kinetic rate constant $k$ on the temperature, $T$

$$
k = A \exp \left( -\frac{E}{RT} \right)
$$

(Eq. 6.4)

$A$ is pre-exponential factor (min$^{-1}$), $E$ is apparent activation energy (J mol$^{-1}$), $T$ is temperature (K) and $R$ is universal gas constant 8.3145 (J mol$^{-1}$ K$^{-1}$). The reaction rate is given by:

$$
rate = kO_2
$$

where rate constant, $k$ is temperature dependent according to the Arrhenius equation and $O_2$ is the concentration of oxygen.

The storage simulated is based on large piles in an indoor storage. Thus, the effect of wind is not considered. Thus, the heat transfer equation used in the convection heat flux on the surface is driven by natural convection. The input parameters determined from the experimental works used in the simulations are shown in Table 6.1. Specific heat and thermal conductivity of both material is taken from literature Koufopanos et al., (1991).

Table 6.1: Model inputs for simulation

<table>
<thead>
<tr>
<th></th>
<th>Non-torrefied sample</th>
<th>Microwave torrefied sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific heat, Cp J/(kg.K)</td>
<td>1112.0+[4.85(T-273)]</td>
<td>1112.0+[4.85(T-273)]</td>
</tr>
<tr>
<td>Thermal conductivity, $\lambda_p$ W/(m*K)</td>
<td>0.13+[0.0003(T-273)]</td>
<td>0.08-0.0001(T-273)</td>
</tr>
<tr>
<td>Bulk density, kg/m$^3$</td>
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<td>500</td>
</tr>
<tr>
<td>True density, $\rho_p$ (kg/m$^3$)</td>
<td>1451</td>
<td>1442</td>
</tr>
<tr>
<td>Porosity, $\varepsilon$</td>
<td>0.695</td>
<td>0.653</td>
</tr>
<tr>
<td>Activation energy, $E_a$ (kJ/mol)</td>
<td>77.37</td>
<td>68.3</td>
</tr>
<tr>
<td>Pre-exponential factor, s$^{-1}$</td>
<td>3.85E3</td>
<td>3.24E4</td>
</tr>
</tbody>
</table>
6.3.2 Mesh

The free triangular shape of mesh with the maximum element size is 0.135 m and the minimum element size is 0.006 m. Element size for mesh chosen for the simulation is the once calibrated for fluid dynamic.

Figure 6.3: Free triangular shape mesh was chosen for the simulation

6.4 Model validation

6.4.1 Introduction

To validate the numerical model discussed in Section 6.3.1 against the experimental data, simulations of the heating behaviour were run based on actual conditions during the study of the thermal runaway in bulk tests. The simulated results were compared to the real data collected during the experiments. Hence, the simulations were based on the effect of the oven temperature towards the bulk sample.

Therefore two separate domains were used to simulate the experiments, namely open domain and porous domain. The porous domain is for the modelling of the wire mesh basket while the open domain modelled the heating up by the oven using air as a conductive medium. The validation model was built in 2-Dimensional axisymmetric. This type of model was chosen to reduce the computation time and modelling time. The validation of the model was done by comparing the temperature profile at the centre of the bulk, which labeled as point T1 in the bulk test. The point T1 is shown in Figure 6.4. The comparison of the centre temperature is chosen because the self-heating process of the sample can only be detected when
temperature at the centre exceeded the oven temperature. Therefore, for validation purposes, only one point will be validated, which is T1. In addition to that, the temperature evaluations that were considered for validation purposes were considered only in the key region, where the results of the time taken for the samples to undergo self-heating that was in good agreement with the simulation and experimental data. Due to the simplification of the model such as eliminating the forced convection from the fan in the oven, the heat up pattern would have some discrepancy compared to the temperature profile from the experiments.

COMSOL Multiphysics® includes the physics-based modules that provide predefined physic interfaces, which can be combined to solve several multi-physics applications. Therefore in this validation, the modules chosen were based on the porous media physic interface, which represented the bulk samples. Therefore the modules used in the simulations are:

(i) Heat transfer in the porous media;
(ii) Species transport in the porous media; and
(iii) Free and porous media flow.

### 6.4.2 Geometry of the model

The mesh chosen for the modelling is the free triangular mesh as well, with the finer mesh at the boundary of the wire basket. The size of the mesh for the whole domain is between 0.0101 m and 0.00045 m. While the size of the mesh at the boundary of the wire mesh were finer, which is between 0.00675 m and 0.0003 m.
6.4.3 Result of the validation

a) Validation of bulks test of microwave torrefied sample

The validation simulation was done for the bulk test using Bulk 1 at 190°C for microwave torrefied sample. The temperature profile in the 2-D axisymmetric geometric is shown in Figure 6.5. The higher temperature is detected at the wire mesh filled with the sample (Bulk 1) after being heated up to 5000 seconds (83.3 minutes). The comparison of the temperature profile is shown in Figure 6.6. The temperature development tends to slow down at the beginning up to 1750 seconds, then exceeded the temperature pattern of the experimental result.

As mentioned earlier, only the key region of the simulation is significant in the model validation. The early stage of the heating process of the simulation did not follow the same pattern as the bulk test. Nevertheless, even though the patterns of the temperature profiles were inconsistent before 1500 seconds, both temperature profiles gave slightly the same induction time, where it can be witnessed that both temperature profiles exceeded oven temperature at almost the same duration. Therefore in Figure 6.6, only the temperature profile after 1500 seconds was discussed.
Chapter 6: Development of numerical models to simulate the self-heating process in bulk storage area

The inconsistent temperature profiles at the beginning of the heating up process for the simulated and the experimented data were predicted due to the simplification of the model simulated. The air convection due to the fan inside the oven was not included in the simulation model. According to Yan et al. (2005), the complexity of the spontaneous ignition problem is due to the strong coupling between the air flow, heat and mass transfer, physical and chemical reaction. Therefore, by deducting one of the significant factors such as airflow due to a fan in the oven, it can prevent the possibility of determining a reliable validation method. Besides that, the loss of moisture during heating process was also ignored in the simulation. These factors were significant reason that would contribute to the inconsistent temperature profile measured for simulation and experimental data. This phenomenon occurred because to the system was trying to balance the heat build up and the heat dissipated with the effect of air circulation in the oven.

![Figure 6.6: The comparison of temperature profiles at the centre for microwave torrefied sample heated at 190°C.](image)

Then the validation was done for the second sample to see whether the pattern is the same or not. The sample chosen is microwave torrefied sample in Bulk 1 heated at 180°C oven temperature. Based on the temperature profiles in Figure 6,7 The temperature patterns of the simulation result follow the one done earlier, with the temperature slightly lower at the beginning and then increased afterwards. However,
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the induction time of the simulation and experimental results showed a good agreement, where they exceeded the oven temperature at the same duration.

Based on Figure 6.6, the temperature at the centre of the simulations results exceeded oven temperature at 3290 seconds. This indicated the self-heating leading to thermal runaway occurred in the sample where the heat increased rapidly after exceeded the oven temperature. However, the time taken was slightly lower than the one determined in bulk test, which is 3513 seconds. However, it can be safe to say that the profile of heating followed the pattern of heating up due to the chemical exothermic reaction in bulk.

In addition to that, the temperature evaluation of the simulation results seems to heat up linearly at the beginning before slowing down a bit afterwards. This phenomenon occurred because the system was trying to balance the heat build up and the heat dissipated with the effect of air circulation in the oven. Thus, in conclusion, the simulation results agreed well with the experiments for the microwave torrefied.

Figure 6.7: The comparison for experimental and simulation for microwave torrefied biomass heated at 180°C in Bulk 1.
b) Validation of bulks test of non-torrefied sample

In order to validate the kinetic parameters of the non-torrefied sample, the model validation was done at an oven temperature of 180°C in Bulk 1. Based on Figure 6.8, the same steady state achieved after the centre temperature reached the oven temperature. The steady state achieved once the sample reached oven temperature, which indicated that there is no self-heating occurred for the non-torrefied sample heating at the specified condition.

![Temperature Comparison](image)

Figure 6.8: The comparison between temperature patterns at the centre for experimental and simulation for non-torrefied biomass heated at 180°C

Based on Figure 6.8, the temperature profile of the simulation results seem to follow the one done in the experiment, but the temperature seems to shift to later time compared to simulation. The temperature pattern is not consistent with the experimental result. However it shows then the same finding, which no self-heat is detected for the non-torrefied sample heated up in Bulk 1 at 180°C. Overall, the simulation results agreed well with the experiments. In particular, the time of induction in the test runs closely matches the model predictions, despite their inconsistent pattern.
6.5 Variable analysis

The spatial distribution of temperature is an important parameter to determine the location of the ignition point. Thus, the results of the variable analysis in this section are examined based on the temperature contours distribution. The variables study are the pile height, pile width and the ambient temperature. The hottest point is the maximum temperature detected within the pile. Thus, the evolution of the hottest point with the ignition induction time for each simulation is used to predict the possibility if the self-heating that lead to ignition within the pile. Based on the spatial distribution, the location of the ignition can be detected. The influences of the variables on the hottest point (maximum temperature) are evaluated.

6.5.1 Effect of pile height

The effect of the pile height is examined for both samples, where the temperature profile presented on the left is for microwave torrefied sample and on the right is for the non-torrefied sample. It was presented side by sides to give an easy comparison of the temperature evolution. To evaluate the effect of the height of pile for both sample, the simulation was done at an ambient temperature of 60°C, at different pile heights of 2, 4, 6, and 8 m for 4500 hours (187.5 days).

The heights are chosen based on the recommendation of stack height in the literature such as (Krigstin & Wetzel, 2016; Hogland & Marques, 2003; Blomqvist & Persson, 2003). Besides that, Murasawa et al. (2012) had proved that it was not practical to pile up the biomass material such as more than 10 meters. Based on their study, the critical ignition temperature of the material can be as low as 50 °C. Therefore, after taking those literature into consideration, those pile heights was chosen for simulation purposes to examine the effect of the pile height on the tendency of the samples to self-heat.
a) Simulation of pile height of 2 m at an ambient temperature of 60°C for 4500 hours.

Figure 6.9: Temperature contours of pile height of 2 m stored at the ambient temperature of 60°C for 4500 hours. Figure 6.9 shows the hottest spots for both samples are at the bottom centre (1.5,0). The highest temperature detected in a non-torrefied sample is lower than the one in microwave torrefied sample. Thus indicated that the microwave torrefied sample is more reactive compared to the non-torrefied sample, where the heat due to the chemical reaction able to dissipated to the surrounding and balances the heat build up in the non-torrefied sample.

The simulation conditions for both samples indicated that no ignition detected after 4500 hours (187.5 days). However, at 4500 hours, the highest temperature detected for microwave torrefied sample was 66.8°C, which greater than the non-torrefied sample which is only 61.8°C. Therefore, 8% increased the highest temperature compared to the ambient temperature of 60°C. Thus, at pile height of 2 m and ambient temperature of 60°C, the storage condition is considered as safe, which no self-heating detected.
b) Simulation of pile height of 4 m at an ambient temperature of 60°C for 4500 hours.

![Temperature contours of pile height of 4 m stored at the ambient temperature of 60°C for 4500 hours.](image)

Based on Figure 6.10, the hottest spot detected for microwave torrefied sample was at coordinate (1.5,2.9). While for non-torrefied sample the hottest spot detected was still at the bottom centre (1.5,0). The highest temperature in the non-torrefied sample is lower that the one in microwave torrefied sample. This indicated that the microwave torrefied sample is more reactive compared to the non-torrefied sample. The heat started to shift to the surface. However, the heat dissipated to the surrounding and balances the temperature build up in the non-torrefied sample. No ignition detected after 4500 hours (187.5 days). However the highest temperature identified is 64.6°C. The highest temperature detected is lower than the one detected in at 2 m pile height storage (as shown in Figure 6.9). The phenomena indicated that the heat generated managed to dissipated to the surrounding and did not accumulate at the bottom of the pile.
c) Simulation of pile height of 6 m at the ambient temperature of 60°C for 4500 hours.

Figure 6.11: Temperature contours of pile height of 6 m stored at the ambient temperature of 60°C for 4500 hours.

Figure 6.11(a) shows the temperature contour of the hottest spot detected for microwave torrefied sample was at coordinate (1.5, 4.9) after 4500 seconds. For a non-torrefied sample, the hottest spot detected is still at the bottom centre (1.5, 0). The highest temperature in the non-torrefied sample is lower than the one in microwave torrefied sample. However, no ignition detected after 4500 hours (187.5 days). The region with highest temperature distribution is shifted to the top of the pile. As the heat losses are tend to be on the surface of the pile.
**d) Simulation of pile height of 8 m at the ambient temperature of 60°C for 4500 hours.**

![Temperature contour of pile height of 8 m stored at the ambient temperature of 60°C for 4500 hours.](image)

Based on Figure 6.12, the hottest spot for microwave torrefied sample is at coordinate (1.5,6.9). While the hottest spot of the non-torrefied sample at 4500 hours is detected at (1.5,3.17). It can be seen that the hot spot started to build up towards the top of the surface at this height for the non-torrefied sample. However, no ignition detected for both samples.

**e) Conclusion of the effect of simulation at various pile heights**

According to the simulations at different heights, we can see that the highest temperature region is detected towards the top surface of the pile. The pattern is observed for all the height simulated. The heat seems to dissipate towards the surface of the pile due to the natural convection set up for the pile. In all cases for the non-torrefied sample, the heat of oxidation is suppressed to the bottom of the pile. The opposite phenomenon is witnessed for the microwave torrefied sample, where the hot spot tends to shift to the surface.
6.5.2 Effect of pile width

The effect of pile width on the self-heating behaviour also being examined using the simulation. Heat due to the exothermic reaction in the system is generated throughout the body, however, only dissipated through the surface. Therefore, considering the width of the stockpile, self-heating leading to ignition can happen even though the stockpile is elevation is not high, especially at the higher ambient temperature. This phenomenon is being simulated to consider the effect of pile width on heat distribution within the pile.

The simulation was performed using microwave torrefied sample stored at an ambient temperature of 80°C with pile size of 5 m (W) x 1 m (H). The simulation was set to run for storage duration of 180 days. However, ignition had been detected at 106.25 days (2550 hours). The temperature contours for the heat distribution at different duration is presented in Figure 6.13.

Based on the Figure 6.13 (a) and Figure 6.13 (b), the temperature distribution of the one at 15 days and 30 days are not much different, with the highest temperature recorded along the surface is 80°C. However, the temperature started to increased rapidly at days 90. The increased of hottest temperature was from 81.7°C for 60 days to 97.3°C for 90°C. At days 60, the temperature development seem to concentrated to the centre with the coldest pile temperature is 67.7°C compared to 29.4°C at 30 days. This indicated that the temperature increase due to exothermic reaction accumulated at the center.

At days 90, the two hottest spot is seen to develop, which indicated that the heat generated at the bottom of the pile trying to dissipate to the surrounding. However, because the distance to the surface is far, the heat distributed to two spots and accumulated instead of dissipated to the surface. Two self-heating spots that lead to ignition can be found in Figure 6.13 (e), where the pile ignited at 106.25 days (2550 hours).
Figure 6.13: Temperature contours of torrefied sample with pile size 1(H) x 5 (W) meter and ambient temperature 80°C.
The temperature profile of the hottest spot (1.5,0) is plotted in Figure 6.14. The exponential rise of the temperature profile can be seen after 2550 hours. The simulation indicated that the temperature profile at the hottest spot shows a supercritical curve. Where the heat production in a pile surpassed the ambient temperature. The thermal runaway is considered to happen here. Even though the pile height is just 1 m, it can be seen here that the heat build up from the exothermic reaction was high as the reaction rate increase with temperature.

![Temperature profile](image)

**Figure 6.14**: Temperature profile at the hottest spot of the microwave torrefied sample after being stored for 2550 hours at ambient temperature of 80°C

### 6.5.3 Effect of ambient temperature

Different ambient temperature can be simulated to examine the effect of ambient temperature on the self-heating tendency of the samples. Therefore a critical ambient temperature can be determined for the samples. The critical ambient temperature is defined as the ambient temperature at which the internal heat generation exactly balances with the heat loss in the system (Morrison & Hart, 2012). Consequently, below are the discussions to examine the effect of ambient temperature on self-heating behaviour.
a) **Relationship of ambient temperature and tendency of the sample to self-heating**

To establish the relationship between the ambient temperature and tendency of the sample to self-heating, simulations were done to observe the heating behaviour of the microwave torrefied sample. The pile size was set to constant of 1 m (W) x 1 m (H) at various ambient temperatures for 180 days. The square shape pile is chosen to eliminate the height effect of the temperature distribution when the pile heated up. Figure 6.15 shows the temperature contour of microwave torrefied sample simulated at various ambient temperatures.

![Temperature contours of microwave torrefied sample heated at various ambient temperatures](image)

Figure 6.15: Temperature contours of microwave torrefied sample heated at various ambient temperatures
Figure 6.15 (a) and Figure 6.15 (b) shows the temperature contours of the pile when the ambient temperature is 40°C and 50°C respectively. Based on the Figure 6.15 (d), the temperature increase is quite small over the pile after 180 days, with just 3.6°C increased after 180 days. Thus the temperature effect of 70°C is still not very significant to started up the heating process due to the chemical reaction. However, if the pile heated up for a longer time, the self-heating will be started to show.

After 180 days., the temperature increased in a pile stored at an ambient temperature of 80°C is 9.3°C. The temperature difference is obvious when the pile stored at an ambient temperature of 80°C. Thus, 80°C is the temperature that can trigger the self-heating leading to thermal runaway. At 85°C, the increased of temperature started to concentrate to the bottom (Figure 6.15 (f)).

The highest temperature achieved when stored at different ambient temperatures were plotted in Figure 6.16. The relationship between ambient temperature and highest temperature reached by the pile can be explained, where at 80°C of ambient temperature, the highest temperature of the pile increased only about 11% from the ambient temperature. However, at 85°C of ambient temperature simulated, the highest pile temperature increased up to 110°C, which is an increase of 29%. Thus, the higher the ambient temperature, the higher the tendency of the pile temperature to increases, which unfortunately will lead to self-ignition.

![Figure 6.16: Relationship between highest pile temperature and the ambient temperature of microwave torrefied sample](image)
Based on the previous study done in the industrial food waste, the material will gradually generate heat through fermentation as well as oxidation process although stored at near room temperature (Murasawa et al., 2012). Therefore in some cases, it can cause self-ignition when the heat generated exceeds the heat that can be dissipated. Furthermore, critical temperature for each material is unique for given surface to volume ratio, which follows the Arrhenius equation as discussed in important findings in self-heating theory by Bowes (1971).

**b) Comparison of the self-heating tendency of both samples**

The effect of temperature was then tested on both samples, and the temperature profiles were presented side by sides to give an easy comparison of the temperature evolution. The pile size is 3 x 8 m, and temperature of 50, 60, 80 and 85 °C were chosen for comparison purposes, and simulation was run for 365 days. The height of 8 meter was chosen in the simulations due to the criticality of the height when simulated in Section 6.5.1. However, not all samples can be simulated up to 365 days as the self-heating lead to ignition occurs before 365 days.

(i) **Ambient temperature of 50°C**

For the first simulation, the ambient temperature of 50°C is chosen. Based on the Figure 6.16, the temperature contours for both samples look slightly the same with maximum temperature increased after 365 days is only 0.1°C.

![Temperature contours at ambient temperature of 50°C](image)

a) Microwave torrefied sample  
(b) Non-torrefied sample  

Figure 6.17: Temperature contours at ambient temperature of 50°C
Therefore, no self-heating was detected at the end of the simulation, which implies that when the sample stored below the critical ambient temperature, the internal heat generation balances with heat loss in the system (Morrison & Hart, 2012). According to the simulations, both samples will theoretically pose no risk of self-heating to ignition if stored at 50°C for a year. However, in order to see the temperature evolution in bulk, the temperature evolution of the hottest point for each sample was plotted in Figure 6.18. A different temperature profile patterns can be seen for the samples, where the non-torrefied samples (Figure 6.18 (b)) shows the steady state curve after 100 days. However, for microwave torrefied sample, a smooth temperature increases can be seen after 150 days. This pattern suggested that there is a possibility for the microwave torrefied sample to undergo self-heating if stored more than 365 days. This simulation stops at 365 days, due to the realism of the storage duration of biomass fuels in energy industry which will not be more than a year. Though there is higher risk of microwave torrefied sample to self-heat than the non-torrefied sample to self-heat, there is still relatively low risk of self-heating occurrence for both samples stored in 50°C.

![Temperature profile of hottest point at ambient temperature of 50°C](image)

(a) Microwave torrefied sample          (b) Non-torrefied sample

Figure 6.18: Temperature profile of hottest point at ambient temperature of 50°C

(ii) Ambient temperature of 60°C

The simulations were extended for the higher ambient temperature of 60°C for 365 days. The results of the temperature contours from the simulations of both samples were shown in Figure 6.19. Based on the temperature contours for both samples, it can be witnessed that the evolution of the hottest point is shifted to the top centre of the pile in the microwaved torrefied sample (see Figure 6.19 (a)). The hottest point is detected at (1.5, 4.5) compared the hottest point of the non-torrefied samples that
occurred at the bottom of the pile. In addition to that, the highest temperature measured in microwave torrefied sample was 145°C compared to only 62.6°C for the non-torrefied sample. The temperature rise in the bulk for microwave torrefied sample indicated that self-heating occurred if stored for 365 days.

![Temperature contour at ambient temperature of 60°C](image)

**Figure 6.19**: Temperature contour at ambient temperature of 60°C

The analysis was extended with the plotted temperature of the hottest spot for both samples. As shown in Figure 6.20, there is an obvious distinctive pattern between Figure 6.20 (a) and Figure 6.20 (b); which in Figure 6.20 (b) the steady state is reached by the non-torrefied sample after roughly 90 days, which is much shorter than the one showed for the non-torrefied sample stored at 50°C. The shorter duration showed that the system achieved steady state faster in higher ambient temperature.

![Temperature profile of hottest point at ambient temperature of 60°C](image)

**Figure 6.20**: Temperature profile of hottest point at ambient temperature of 60°C
However, for microwave torrefied sample, a supercritical behaviour is detected. However, no ignition is detected for the non-torrefied samples stored at 60°C; it is unlikely that this condition simulated would have reached spontaneous ignition witnessed in Figure 6.20 (a). The rapid increase of temperature can be seen and the thermal runaway pattern was detected at 320 days. The same supercritical behaviour that indicated the self-ignition due to self-heating process can also be seen in the several studies of self-heating behaviour (Chen, Sidhu, & Nelson, 2013; Ramírez, García-torrent, & Tascón, 2010; Jones & Vais, 1991). The comparison of the hottest point in Figure 6.20, shows that the non-torrefied sample is much less reactive in both temperatures, however at the ambient temperature of 60°C the reaction rate increased. The exothermic reaction in of the system seems to increase with the increase of the ambient temperature (Garcia Torrent et al., 2015). Therefore the steady state is reached earlier when simulated in an ambient temperature of 60°C compared to the one simulated in an ambient temperature of 50°C.

(iii) Ambient temperature of 80°C

To examine the effect of higher ambient temperature, 80°C was chosen for simulations. Based on Figure 6.21, a different pattern of the temperature contours of the pile at an ambient temperature of 80°C is witnessed. The hottest region of the microwave torrefied sample followed the same pattern as observed at an ambient temperature of 80°C, where the location of the hottest region shifted to the surface of the pile.

Figure 6.21: Temperature contour of hottest point at ambient temperature of 80°C
The hottest point was increased to 156°C from the ambient temperature simulated and the hottest point is detected at (1.5, 7.0). Also, the non-torrefied sample, the hottest point is shifted to the upper part of the pile, which is at (1.5, 2.5).

The analysis was extended with the plotted temperature profiles in Figure 6.22, which shows the temperature evolution of the hottest spot for both samples. There were thermal runaway phenomena detected for both samples, which the supercritical curves can be witnessed for both samples, even though the point of exponential increased is much longer to reach for the non-torrefied sample. The comparison of the temperature profiles showed that microwave torrefied sample achieved a supercritical stage of rapid temperature rise, which started at 130 days.

Therefore, for the storage are with an ambient temperature of 80°C, it is safer to store the non-torrefied sample up until 170 days, compared to the microwave torrefied sample, which reached the rapid heating stage at 130 days only. The results achieved from the simulations were in good agreement to the reactivity determined for both samples. Therefore the patterns follow the finding that the microwave torrefied samples are more reactive compared to the non-torrefied sample during storage.

(iv) Ambient temperature of 85°C

Lastly, the simulation was extended to the ambient temperature of 85°C. Figure 6.23 shows the temperature contours of the pile, where obvious hottest spot can seem for both samples, regardless of the locations. The highest temperature recorded before the thermal runaway occurred was 153°C for both samples.
Chapter 6: Development of numerical models to simulate the self-heating process in bulk storage area

As shown in Figure 6.24, both samples took shorter time to ignite compared to the one simulated in 80°C. At 85°C, the microwave torrefied sample started to ignite at 111 days as illustrated in Figure 6.24 (a), while the sample began to ignited at 120 days for the non-torrefied sample (Figure 6.24 (b)). It can be safely concluded that a safe storage practice can be achieved when stored below 80°C because a rapid temperatures increase for both samples were detected in both samples. The exponential increased of the temperature was double its ambient temperature.

Therefore, based on the results discussed earlier, the safe storage condition for such pile can be simulated. In order to establish the relationship between the ambient temperature and time taken for the pile to achieve its thermal runaway is shown in Figure 6.25.
Based on the relationship between the ambient temperature and induction time for microwave torrefied sample, if the ambient temperature reached 80°C, the microwave torrefied sample cannot be kept more than 133 days. However, based on this thermal stability map, for long term storage, the microwave torrefied sample is safer to be stored at normal room temperature, or somehow below 50°C. Therefore microwave torrefied material is safe to store in the short-term period, for example, less than three months to avoid the self-heating that lead to self-ignition of the samples to occur.

6.6 Discussions on effect of the variables

Morrison & Hart (2012) had stated that the possibilities for a biomass bulk material to self-heat is an important chemical reactivity hazard that must be effectively managed to limit the possibilities of thermal runaway to occur due to self-heat. The result of the effect of height towards self-heating tendency showed the same conclusion as made by researchers such as Zhu et al. (2013) on coarse coal stockpile, Jirjis (2005) on willow shoots, Guo (2013b) on wood pellets. All the researchers had concluded that effect of height could be an ideally controllable variable to avoid the spontaneous heating problem from occurring.
Based on the finding from Section 6.5.1, we can see that the shorter pile height will make it easier for heat accumulated in the system to dissipate to the surrounding through the surface area. All the increase of the temperature determined from the simulation is less than 7% increase. Furthermore, even though no ignition detected when the samples simulated for each of the simulation, it is still a bad practice to let a biomass pile heat, as it will degrade the quality of the biomass itself.

Storage below critical ambient temperature leads to a steady state condition, which the heat dissipated to the surrounding balances the internal heat generation. Morrison & Hart (2012) described that the biomass will theoretically pose no risk of self-heating that leads to self-ignition. Section 6.5.2 showed that, when the ambient temperature increases, the temperature of the pile gradually increased towards a rapid temperature increased. Consequently, a critical ambient temperature should be determined to achieve a safe storage and handling condition. Many researchers such as Luo et al. (2016); Larsson et al. (2012); Bygbjerg & Pauner (2006) managed to find the unique relationship between critical ambient temperature and the type of the biomass tested as well as the size of the pile. The higher critical ambient temperature is determined for the non-torrefied sample due to its reactivity. Based on the reactivity study done, it is concluded that the microwave torrefied sample is more reactive, thus the critical ambient temperature for handling and storage purposes is lower compared the non-torrefied sample.

The biomass fuels need to be store in a smaller pile, to avoid the self-heating that lead to ignition. In addition to that, existing practice in the biomass handling and storage also prevent the pile of the stockpile in great width. Therefore, for practicality and safety issues, the pile should be piled up horizontally and avoid a long horizontal pile. This conclusion is in agreement with the findings by Saddawi et al. (2013).
6.7 Summary

- The validation of simulation model agreed well with the experiments. In particular, the time of induction in the test runs closely matches the model predictions, despite their inconsistency in temperature pattern. The highest temperature region tends to fall at the surface at the pile.

- At 80°C of ambient temperature, the maximum temperature of the pile increased about 11% from the ambient temperature and at 85°C of ambient temperature simulated, the highest pile temperature increased up to 110°C, which is an increase of 29%.

- The exponential graph is plotted based on the relationship between the ambient temperature and the highest temperature within the piles. The exponential increase is detected after the ambient temperature of 80°C.

- The simulation can provide some suggestions on how to control the parameters to retard or suppress the self-heating taking place in the biomass fuel piles and as the guides in performing a larger scale of simulations.

- Based on the simulation, it can be concluded that the microwave torrefied sample is more reactive than the non-torrefied sample.

- 80°C is considered as critical ambient temperature for the storage condition of the pile, where the exponential pattern is observed to start at 80°C for 1x1 m pile of microwave torrefied sample in the relationship between highest pile temperature and the ambient temperature.

- The height and ambient temperature are the critical parameters in the storage of bulk biomass and the higher ambient temperature, the faster the pile ignited.
Chapter 7 Conclusions and Recommendations for Future Research

7.1 Conclusions

In this work, the self-heating behaviour of the non-torrefied and microwave torrefied biomass has been studied experimentally and numerically. The conclusions from the study are:

- The study has shown that the microwave torrefaction technology can improve the combustion properties of the biomass fuel. The properties of torrefied biomass are comparable to the one in low ranking coal. The calorific value of torrefied biomass increased to 22MJ/kg from 18.4MJ/kg in the non-torrefied sample. The O/C and H/C ratio decreased after the torrefaction process, where the O/C ratio is 0.67 in the non-torrefied sample and 0.45 in microwave torrefied sample. The H/C ratio is 0.14 in the non-torrefied sample and 0.11 in microwave torrefied sample.

- The proximate analysis showed that the volatile matter in biomass was reduced from 85.6% to 68.7% after the torrefaction. While the fixed carbon content is increased from 3.9% to 18.6% after torrefaction. The ash content in torrefied biomass samples rose to 10.5% from only 4.7% in the non-torrefied sample.

- The thermal degradation was carried out using thermogravimetric analysis for both torrefied and non-torrefied sample in air and nitrogen with the heating rate of 5°C/min to obtain the temperature of maximum weight loss and the temperature of the initial combustion.

- Thermogravimetric analysis in the air has shown that the temperature maximum weight loss (TMWL) for the non-torrefied sample is 310°C, while for the torrefied sample is 312°C. The temperature of the initial combustion (TIC) 268°C for non-torrefied sample 283°C for microwave torrefied sample.
• Thermogravimetric analysis in oxygen has shown that the characteristic temperature ($T_{\text{charac}}$) for non-torrefied sample is higher than the torrefied sample. $T_{\text{charac}}$ non-torrefied sample, which is 331°C and 329°C respectively.

• The activation energy derived from thermal decomposition in air of the non-torrefied sample is between 64.5 to 84.4 kJ/mol, while for torrefied sample the range is lower which is between 53.1 to 70.7 kJ/mol. The activation energy for thermal decomposition in air shows that microwave torrefied sample is more reactive comparing to non-torrefied sample.

• The lower activation energy for torrefied biomass fuel indicated that the material is reactive and have a higher tendency to self-heating. The risk ranking graph; is drawn based on the activation energy and characteristic temperature of the samples against other materials from literature. Both samples are classified as a medium risk, but with microwave torrefied sample in the rank of higher tendency to self-ignite comparing to the non-torrefied sample.

• The test showed that when the oven temperature is below and at 180°C no ignition is detected for the non-torrefied sample. However, microwave torrefied sample started to self-heat at 4002 seconds (66.7 minutes). It can be concluded that the torrefied sample is more reactive compared to the non-torrefied sample.

• Series of the bulk test is carried out in the uniformly heated oven at constant air temperature to study the thermal heating behaviour independently. The test showed the ignition induction time for microwave torrefied sample heated at 180°C is 4002 seconds at Bulk 1, 5066 for Bulk 2 and 7611 for Bulk 3. The ignition induction time increased with the decreasing of the bulk volume. The ignition induction time is decreasing when the oven temperature is increased.
The numerical simulation of heat and mass transfer was modelled using COMSOL Multiphysics® modules, consists of heat transfer in the porous media, species transport in the porous media as well as free and porous media flow using the kinetic parameters from the experimental data to determine the heat generated from the exothermic reaction.

The simulations carried out for the bulk tests for both samples. The temperature profiles obtained in the simulation were compared with the experimental data, and it was shown the temperature profiles in good agreement between measured and simulated. Therefore, the numerical model is validated against the experiments and could be used for the study of other conditions in confidence.

Based on the simulation, the safe practice to store the biomass is by piling up horizontally is safer than horizontally. The resulting form the simulations show two hot spots can be detected when the pile was pile with a wider width.

80°C is considered as critical ambient temperature for the pile, where the exponential pattern is observed to start at 80°C in the relationship between highest pile temperature and the ambient temperature of the microwave torrefied sample in 1x2 m pile.

Examples of simulation in large storage area presented here showed that the model could predict the development of the hot spot in the piles, therefore can be used to help identify the possible self-heating phenomena that lead to ignition.
7.2 Recommendations for future research

Although the self-heating propensity of biomass material had been extensively studied, more research is needed for the biomass fuels such as torrefied material especially the fuels produced using microwave torrefaction method. This works managed to show that the higher reactivity of fuels produced using microwave torrefaction method. The methods of experimental study and numerical model developed can be used for other types of biomass material, as well as simulations with different storage conditions. Therefore, it would be interesting to understand the effect of torrefaction process parameters such as reaction time and reaction temperature towards self-heating parameters. In addition to that, the effect of the various torrefaction methods on self-heating propensity is an interesting topic to focus on too. The methods of torrefaction that produced a material with the less reactive fuels could be determined. Furthermore, it would be interesting to study the effect of physical properties of the torrefied biomass fuels such as porosity, density and moisture content towards its self-heating propensity. Therefore a safe handling of the fuels with upgraded physical properties from the torrefaction process can be achieved.

Finally, the simulation works on the storage of biomass fuels can also be extended towards the effect of the physical properties of the fuels. For example, the relationship between the physical properties and the critical storage condition can be established. Besides that, various storage conditions can also be studied, which could include the possibilities of outdoor storage with the study of critical wind velocity. Furthermore, the geometry of the piles could be simulated in another shape such as the pile with side slope. The works can also be extended to the simulation of the storage condition with different oxygen concentration as well as the possibilities of inert storage condition.
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Appendix A: Thermocouples readings for bulk tests

a) 

b) 

c) 

d) 

e) 

Appendix A - 1: Thermocouples readings of microwave torrefied sample heated in 180°C for Bulk 1
Appendix A - 2: Thermocouples readings of non-torrefied at 200°C in Bulk 1
Appendix A – 3: Thermocouples readings of non-torrefied sample at 200°C in Bulk 2
Appendix A – 4: Thermocouples readings of non-torrefied sample at 200°C in Bulk 3
Appendix A – 5: Thermocouples readings of microwave torrefied sample heated at 180°C in Bulk 1
Appendix A

Appendix A - 6: Thermocouples readings of microwave torrefied heated at 180°C in Bulk 2
Appendix A - 7: Thermocouples readings of microwave torrefied sample at 180ºC in Bulk 3
Appendix A - 8: Thermocouples readings of microwave torrefied sample at 170°C in Bulk 1
Appendix A – 9: Temperature profile for microwave torrefied sample at 190°C in Bulk 1
Appendix B: Publication originated from the thesis