Radiant Ignition of Polyurethane Foam: The Effect of Sample Size

Rory Hadden, Abubakar Alkatib, Guillermo Rein and José L. Torero, School of Engineering, University of Edinburgh, Edinburgh, UK

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Abstract. Although smouldering ignition of upholstery items remains a leading cause of residential fire deaths, relatively little research is conducted on the topic. An experimental investigation of the effect of sample size on the ignition and spread of smouldering and flaming in polyurethane foam under natural flow conditions is reported here. Polyurethane foam samples are used because this is a common material in modern, residential environments and one for which there exists significant quantities of previous experimental data in the literature. Samples of different square cross-section size and a fixed height of 150 mm are insulated on all sides except the top which is exposed to a radiant heat flux and is open to the air. Samples with side lengths of 50 mm, 100 mm, and 140 mm are studied. Ignition and spread dynamics are diagnosed using thirteen thermocouples located along the vertical centre line. The onset of smouldering ignition (13 kW m\(^{-2}\), 8 kW m\(^{-2}\) and 7 kW m\(^{-2}\) for 50 mm, 100 mm and 140 mm sample sizes respectively) is observed at significantly lower heat fluxes than flaming (45 kW m\(^{-2}\), 32 kW m\(^{-2}\) and 30 kW m\(^{-2}\) respectively). Critical heat fluxes for smouldering and flaming ignition increase with decreasing sample size, with smouldering ignition being significantly more sensitive to sample size than flaming ignition under the size range studied. Smouldering spread rates are measured in the range from 3 mm min\(^{-1}\) to 25 mm min\(^{-1}\) and found to be a strong function of the heat flux and depth of the smoulder front. The effect of sample size on smouldering has been theoretically proposed before but this is the first time that this effect has been demonstrated experimentally for ignition. The fact that large samples result in the lowest critical heat flux could have implications for testing procedures and translation of results from small-scale testing to real-scale in the built environment.

Keywords: Ignition, Sample size, Smouldering, Transition to flaming, Cone calorimeter, Smoulder propagation

1. Introduction

Previous work on smouldering ignition of polyurethane (PU) foam has focused on the effects of flow through the porous fuel [1–3], enhanced or reduced oxygen concentration [4] and the heat flux and exposure time [5]. In this work we present the
effect of sample size on the ignition and spread of smouldering under atmospheric oxygen concentrations and natural flow conditions.

Smouldering is a slow, low-temperature, heterogeneous form of combustion in which oxygen directly attacks the surface of a solid fuel [6–8]. Ignition is governed by the balance between heat flux applied to the surface, heat losses and fuel kinetics [6, 8], while the self-sustained propagation is governed primarily by the oxygen supply to, and heat losses from the reaction front [8–10]. The heat losses are a function of surface area of the front, and the heat released from the reaction depends on the volume of the reaction front; therefore, smouldering combustion is sensitive to the size of the sample [1, 2, 8]. This work aims to experimentally identify the effect of size on the ignition and subsequent propagation of smouldering and flaming in PU foam over a wide range of heat fluxes using three sample sizes.

The effect of sample size on the smouldering ignition of dusts has been studied by Krause and Schmidt [11, 12] and Schmidt et al. [13]. These investigations used hot particles [11] or elevated ambient temperatures [12, 13] to initiate the smouldering reaction. Ignition of larger sample sizes is found to require lower critical ambient temperatures than smaller sizes due to the reduced heat losses from the reaction front. However, larger sample sizes result in lower temperatures and slower propagation of the smouldering front during the reaction. This is attributed to lower rates of oxygen diffusion through the sample to the reaction zone. Schartel et al. [14, 15] studied the effect of sample thickness on flaming ignition of polymers and show that the ignition delay time is reduced and peak heat release rate is increased for thermally thin samples compared to thick samples. These studies did not address smouldering ignition nor the effect of sample area. Ritchie et al. [16] studied the effect of sample area on the heat release from flaming wood. They found that, other than the initial peak, the heat release rate is nearly independent of the sample area. They did not study the effect of size on the critical heat flux for ignition nor smouldering combustion.

The theoretical critical minimum size for sustained smoulder propagation in which the heat losses are balanced by heat generation under conditions of forced flow and no gravity is given by Equation 1 [8],

\[
L_c = \frac{4\delta}{Q_{sml}} \frac{U_{loss} (T_{sml} - T_0)}{\dot{m}''},
\]

where \(L_c\) is the critical length of the sample (side length for a square cross section or diameter of a circular cross section), \(\delta\) is the thickness of the smouldering front, \(Q_{sml}\) is the heat of reaction by smouldering, \(U_{loss}\) is the global heat loss coefficient, \(T_{sml}\) is the smouldering temperature, \(T_0\) is the ambient temperature and \(\dot{m}''\) is the mass flux of oxidizer to the reaction zone.

Using typical experimental data for smouldering PU foam with forced oxidizer flow (\(\delta = 0.04\) m, \(Q_{sml} = 5880\) kJ/kg-O\(_2\), \(U_{loss} = 14\) W m\(^{-1}\) K\(^{-1}\), \(T_{sml} = 450\)°C and \(T_0 = 20\)°C) [1, 2], a minimum sample size on the order of 160 mm is found to be required to sustain smouldering propagation [8]. Equation 1 is only valid for forced flow through the PU foam; however, the scenarios of most importance in fire safety occur under natural flow conditions. In these situations, Equation 1
cannot be applied and it is necessary to study experimentally the critical size problem under natural flow conditions.

Polyurethane foam can undergo smouldering and flaming combustion. At low heat fluxes, a smouldering reaction will be initiated and will propagate through the sample; at higher heat fluxes, flaming ignition will occur. There are two possible mechanisms which may lead to flaming: either transition from a smouldering reaction or auto-ignition in the gas phase. Transition is a complex phenomenon which, though significant in fire safety, has received relatively little study. Putzeys et al. [17] found that the smoulder velocity and peak temperature are strongly correlated to the occurrence of transition from smouldering to flaming. The transition has also been observed to occur when pores are formed in the region behind the smouldering front [18]. This implies that the smouldering front acts as the source of combustible vapours and heat for the flaming ignition. As a result, the onset of flaming ignition may also be affected by the sample size. Flaming ignition may also occur due to spontaneous ignition of the fuel vapours. This requires that the mixture of vapours and air is heated to a temperature sufficient to initiate a gas-phase reaction [7].

2. Experimental Set-Up

The experimental set-up used for these experiments (shown in Figure 1) was based on that of Anderson et al. [5], and not dissimilar to that used by Gratkowski et al. [19]. The set-up is deliberately based on the cone calorimeter, except that there is no gas analysis and the sample depth is increased to 150 mm. A calibrated cone heater provided a uniform heat flux up to 50 kW m$^{-2}$ over the free surface of the sample. The sample was held in a stainless steel sample holder which was inlaid with insulation board to minimize heat losses as well as prevent ingress of...
oxygen from any face other than the heated free surface. The bottom of the sample holder was not inlaid with insulation. As a result, oxygen could only diffuse downward from the free surface. In order to minimize the formation of recirculation eddies induced by the buoyant flows generated by the heater, a 100 mm wide flange was added to the top of the sample holder.

Samples of PU foam with square cross sections of length 50 mm, 100 mm and 140 mm were tested. The depth of all samples was 150 mm. The largest sample was 140 mm long as this is maximum size that can fit under the cone heater and still be subject to a uniform heat flux across the entire surface. The total error in the heat flux measurement was calculated to be ±0.5 kW m⁻², due to the positioning of the heat flux meter and the sample being accurate within ±5 mm. This is in addition to a 3% uncertainty of the meter calibration.

Heat flux along the radius from the sample centre was measured and a maximum reduction of 30% of the centreline value was found at the edge of the 140 mm sample. However, because ignition occurs at the centre of the sample where the heat flux is greatest and heat losses are minimum, this variation in heat flux was not expected to have a significant impact on the results. The vertical separation between the heater and the sample was fixed at 25 mm.

The foam samples were instrumented with thirteen sheathed K-type thermocouples of 1.5 mm diameter. These entered the sample perpendicular to the applied heat flux and were located on the central axis, starting 5 mm below the exposed surface, and subsequently at 10 mm intervals to a depth of 125 mm. Thermocouples were held in place using a vertical stand with holes at locations corresponding to the measurement points on the sample. Temperature data were recorded every second using an Agilent 34980A Multifunction Switch interfacing with a computer running Matlab 2008b. Visual observations and temperature measurements allowed the differentiation between no ignition, smouldering ignition and flaming ignition.

A commercially available PU foam sourced from the UK was used. The density was in the range 20–22 kg m⁻³. The foam is the same as that used by Bustamante et al. [20] who found the elemental composition to be 61.9% carbon, 22.5% oxygen, 8.5% hydrogen and 5.9% nitrogen with trace quantities of sulphur and chlorine. Based on this composition, the overall chemical formula is CH₁.₅₃O₀.₂₇N₀.₀₈.

Prepared samples were placed under the heater and initially shielded from the heat flux using fibreboard insulation. When the heater had reached the temperature corresponding to the desired heat flux at the sample surface, the fibreboard was rapidly removed exposing the sample to the heat flux. No pilot flame or sparking device was used in the set-up.

The critical heat flux required for smouldering and flaming ignitions was found using a systematic, iterative, bisection method. The samples were exposed to a heat flux and, if (flaming or smouldering) ignition occurred, a lower heat flux was selected and a new sample tested. If no ignition occurred, a higher heat flux was used. This method allows the fastest determination of the critical heat flux for ignition. Once the critical heat fluxes for flaming and smouldering ignition had been found, the ignition behaviour at heat fluxes between the two critical heat fluxes and up to 50 kW m⁻² was studied. The number of experiments carried out
for each heat flux and sample size are shown in Figure 2. The experiment was terminated if no ignition was observed after 30 min or when the thermocouple traces reached steady state conditions.

3. Results

Figure 3 shows samples before and after testing. As the heat flux is increased and the combustion becomes stronger, the samples become increasingly destroyed. These images provide qualitative evidence of the combustion processes taking place which is augmented by detailed temperature measurements.

Temperature data measured in the samples are used to distinguish the different types of ignition upon exposure of the sample to a heat flux. The types of ignition are classified as follows: no ignition, partial smoulder, complete smoulder and flaming ignition. Peak temperatures for smouldering were typically in the range 300°C to 400°C, while peak temperatures measured during flaming were 600°C to 700°C, except in the case of the 50 mm samples when flaming was visually

Figure 2. Test matrix showing the heat fluxes at which each sample size was tested. The one-digit numbers indicate how often each experiment was undertaken.

Figure 3. Images of 100 mm samples showing (a) virgin foam, (b) charred foam in which a smoulder front did not propagate, (c) a sample in which smouldering occurred and (d) a sample which underwent flaming ignition.
observed but temperatures in the sample remained around 350°C to 450°C. After the combustion was complete, the thermocouples were allowed to reach steady state before the experiment was terminated.

The reference temperature profiles measured in the apparatus for 100 mm size set-up without a foam sample inside at heat fluxes of 14 kW m⁻², 28 kW m⁻², 44 kW m⁻² and 56 kW m⁻² are shown in Figure 4a. As the heat flux is increased, the steady state temperatures reached by the thermocouples increases linearly. Temperatures are uniform up to a depth of 65 mm after which the temperature begins to decrease slightly. Figure 4b shows that mean steady state reference temperature scales linearly with applied heat flux. Error bars indicate the maximum and minimum recorded temperatures. If the thermocouples exceed the steady state temperatures given by this analysis during an experiment, it can be assumed that there is a significant exothermic reaction occurring. Therefore, these are presented as a reference to identify in the temperature profiles when and where combustion is occurring. Note that temperatures behind the combustion front may remain high because the thermocouples are heated by the smouldering front and insulated by the residue char, which prevents them from cooling to ambient temperature.

The temperature data discussed in the following sections are presented in two ways. The first is an analysis of the temperature–time series, showing the temperature evolution at each thermocouple in the sample over the time of the experiment. This allows the progress of the combustion reaction over time to be assessed. The second is an analysis of the temperature profile through the sample depth at three or four different times during the reaction. These times are chosen to be just after exposure to the heat flux, during combustion and during the final steady state period. This method allows us to see the propagation of a heat wave through the sample. The error in the measurements is given as the mean error for all experiments where repeats were carried out. This was calculated to be 17% for the temperatures.
3.1. 50 mm Samples

Figure 5 shows temperature data for four experiments at heat fluxes 10 kW m\(^{-2}\), 18 kW m\(^{-2}\), 37 kW m\(^{-2}\) and 39 kW m\(^{-2}\). After 40 min, exposure to a heat flux of 10 kW m\(^{-2}\) did not lead to ignition of the foam and only in-depth conduction was observed with no exothermic or endothermic reactions taking place (Figure 5a). The temperature profile shown in Figure 5b shows that the temperature throughout the sample increases in time with the maximum temperature always observed at the free surface. The temperature decreases with depth, as expected in an inert solid. The maximum observed temperature was 275°C.

A heat flux of 18 kW m\(^{-2}\) (Figure 5c and 5d) shows the onset of smouldering at 3 min to 4 min. This is shown in the temperature profiles as an increase in temperature at depths of 25 mm and 35 mm at 5 min compared to the temperature at 15 mm. The maximum temperature is 337°C.

A stronger smouldering reaction was observed upon exposure to a heat flux of 37 kW m\(^{-2}\) (Figure 5e). The temperature profile Figure 5f shows that after 5 min of exposure, the front has spread to a depth of 65 mm and shows a significant temperature gradient between 65 mm and 85 mm. The maximum observed temperature was 461°C.

Figure 5g shows an experiment where a higher heat flux (39 kW m\(^{-2}\)) resulted in flaming and more extensive consumption of the foam, allowing the thermocouples above the flame to measure the temperature of the hot gases (up to 700°C). The flaming did not propagate through the sample and the reaction was quenched at around 105 mm (Figure 5h). The temperature profile shows that there is rapid propagation through the sample and after 1 min, heat has reached the thermocouple 55 mm from the free surface.

3.2. 100 mm Samples

Figure 6 shows temperature traces for experiments at four heat fluxes 8 kW m\(^{-2}\), 8.3 kW m\(^{-2}\), 32.5 kW m\(^{-2}\) and 33 kW m\(^{-2}\). No ignition was observed at a heat flux of 8 kW m\(^{-2}\) (Figure 6a). The peak in the thermocouple near the surface at about 4 min (298°C) was due to degradation of the foam at the surface and exposure of the thermocouple directly to the radiant heat. This is observed in Figure 6b where at 15 min the peak temperature is seen at the thermocouple 15 mm below the surface.

Partial smoulder ignition at a heat flux of 8.3 kW m\(^{-2}\) propagated approximately 55 mm into the sample after 20 min of heating (Figure 6c). Temperatures exceed 300°C for a significant time as the smoulder front propagates through the sample. The maximum temperature is 377°C at a depth of 15 mm. The propagation can be seen in Figure 6d as the peak temperature moved from 15 mm at 8 min to 25 mm after 15 min.

A heat flux of 32.5 kW m\(^{-2}\) (Figure 6e) results in complete smoulder of the sample enhanced by the higher heat flux. The temperature behind the smouldering front is 250°C to 305°C, compared to 75°C to 275°C in Figure 6c. The front propagated from 15 mm to 65 mm between 1 min and 3 min after exposure. The final steady state temperatures are around 300°C as shown in Figure 6f.
Figure 5. Temperature traces and vertical profiles for selected experiments on 50 mm sample (a, b) 10 kW m\(^{-2}\), (c, d) 18 kW m\(^{-2}\), (e, f) 37.3 kW m\(^{-2}\), (g, h) 39 kW m\(^{-2}\). In the left column, the leftmost line corresponds to temperature 5 mm beneath the surface. Moving right, the lines represent depths increasing by 10 mm up to 125 mm.
Figure 6. Temperature traces and vertical profiles for selected experiments using 100 mm sample (a, b) 8 kW m$^{-2}$, (c, d) 8.3 kW m$^{-2}$, (e, f) 32.5 kW m$^{-2}$, (g, h) 34.6 kW m$^{-2}$. Distance form the surface increases from left to right. In the left column, the leftmost line is the temperature at the thermocouple 5 mm below the surface increasing in depth at 10 mm intervals to 125 mm.
Flaming ignition is seen at a heat flux of 34.6 kW m\(^{-2}\) (Figure 6g), where the maximum temperature is 715°C. Figure 6h shows the rapid spread through the foam and steady state temperatures lower than those observed for the complete smoulder case. This is because the foam surrounding the thermocouples has been consumed by the flame, exposing them to increased heat losses. After flaming ignition, the front propagated through the sample in 2.5 min compared to 20 min for smouldering.

### 3.3. 140 mm Samples

Figure 7a and 7b show an experiment at a heat flux of 7.3 kW m\(^{-2}\) in which no smouldering occurred. Smouldering is observed at 7.8 kW m\(^{-2}\) (Figure 7c) with ignition occurring after 10 min exposure. The maximum temperature was 398°C. The smoulder front propagated to 55 mm at 22 min after exposure as shown in Figure 7d.

At 28.8 kW m\(^{-2}\), complete smoulder of the sample is observed (Figure 7e) reaching a maximum temperature of 383°C. The temperature profile Figure 7f shows propagation of the smoulder front with steep temperature gradients over short distances indicating the position of the hot oxidation and colder pyrolysis fronts. Steady-state temperatures of 275°C to 375°C are observed.

Flaming ignition is shown in Figure 7g for a heat flux of 30.6 kW m\(^{-2}\). Rapid propagation is observed in Figure 7h with the front moving from a depth of 35 mm to 85 mm in 30 s. A maximum temperature of 711°C is reached. As in the 100 mm samples, the thermocouples reach a lower steady state temperature than in the smouldering cases.

### 3.4. Summary of Ignition Tests

Figure 8 shows the number of times at which each sample size was tested at a given heat flux. The shading represents the ignition type with light gray representing no ignition, mid gray representing smouldering ignition and dark gray flaming ignition. We can clearly see that the flaming ignition of the 50 mm samples is more scattered than the larger samples. We can also see that there is a significant decrease in the critical heat fluxes required for both smouldering and flaming ignition as the sample size is increased. The numbers indicate how many experiments were undertaken at a given heat flux.

### 3.5. Critical Heat Flux for Ignition

Smouldering and flaming ignitions for the three sample sizes were observed visually as well as using the temperature data seen in Figures 5, 6, and 7. The critical heat flux ranges are detailed in Table 1. These results are in agreement with the work of Anderson et al. [5] and Bustamante et al. [20] where comparison is possible, and complement them with new findings and a wider range of conditions.

The trend observed is that as sample size is increased, the critical heat flux for smouldering and flaming ignition is reduced. This is in agreement with the theory that smouldering is significantly influenced by heat losses from the sample sides.
Figure 7. Temperature traces and vertical profiles for selected experiment using 140 mm samples (a, b) 7.3 kW/m², (c, d) 7.8 kW/m², (e, f) 28.8 kW/m², (g, h) 30.6 kW/m². In the left column, distance form the surface increases from left to right. The leftmost line is the temperature at the thermocouple 5 mm below the surface increasing in depth at 10 mm intervals to 125 mm.
and scales with the surface area to volume ratio [8]. The dependence on heat flux seems to be reduced at sizes greater than 140 mm suggesting that the critical heat flux becomes independent of sample size for large samples.

### 3.6. Maximum Temperature and Time to Ignition

Figure 9 shows the maximum temperatures observed in each experiment. The figure shows three regimes: the first, at low heat fluxes when ignition does not occur, shows a steady increase in maximum temperature with heat flux up to around 300°C. These cases of no ignition are marked by crosses. As the heat flux is increased, a step in the maximum temperature up to around 400°C is observed. This region is represented by circles and corresponds to smouldering ignitions. This step is more pronounced for the 100 mm and 140 mm samples than the 50 mm samples, suggesting a different behaviour for the onset of smouldering at this size. The maximum temperature then increases with heat flux up to approximately

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**Figure 8.** Test matrix showing the heat flux and sample size relationships tested and the resulting ignition types. **One-digit numbers** indicate how many times samples were tested at the heat flux.

**Table 1**  
**Critical Heat Flux for Ignition of the Sample Sizes Studied and Those Found in the Literature**

| Sample size (mm) | Critical heat flux for smouldering ignition (kW·m⁻²) | Critical heat flux for flaming ignition (kW·m⁻²) |
|------------------|-----------------------------------------------------|-----------------------------------------------|
| 50               | 18–19                                               | 32–45                                         |
| 100              | 8–9                                                 | 32–37                                         |
| 140              | 7–8                                                 | 30–31                                         |
| 100 [20]         | N/A                                                 | 35                                            |
| 300 [5]          | 6                                                   | N/A                                           |

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425°C before a second step to temperatures in the region of 700°C. This corresponds to flaming ignition and is denoted by triangles. The critical heat fluxes found by this analysis are in the ranges in Table 1. This diagnostic validates the definition of 350°C as the ignition criterion for smouldering.

The time to ignition for experiments in which smouldering or flaming occurred is shown in Figure 10. The time to ignition was defined when the thermocouple located 25 mm from the free surface reached 350°C. This condition was defined a posteriori based on the analysis presented in Figure 9. For smouldering ignitions, time to ignition ranged from 2 min to 20 min, whereas for flaming this was less than 2 min and flames were generally visually observed less than 5 s after exposure to the heat flux.

Figure 10a shows the time to ignition as a function of incident heat flux. It is observed that the time to ignition decreases as the heat flux is increased. For smouldering initiation, at high heat fluxes, the time to ignition decreases significantly to less than 30 s. It is seen that the ignition time of the 50 mm samples follows a different trend from that of the 100 mm and 140 mm samples.

Figure 10b shows the inverse of the square root of time to ignition. All samples follow a linear relationship as predicted by classical ignition theory [21] indicating that as the applied heat flux is increased, the time to ignition decreases. This suggests that the processes are driven by heat conduction. Again, the 50 mm samples appear to follow a different trend from the 100 mm and 140 mm samples suggest-
ing that the controlling mechanisms of ignition at that size are different from those for larger samples.

3.7. Smouldering Spread Rate

Spread rate was found to be a function of the applied heat flux as shown in Figure 11. Spread rate was calculated between the thermocouples located 25 mm and 35 mm below the free surface. This location is chosen because the heat directly transferred from external heat flux is lower than for locations closer to the

Figure 10. Time to ignition showing (a) asymptotic behaviour and (b) $1/\sqrt{t_{ig}}$ for smouldering (circles) and flaming (triangles) samples. Red, green and blue symbols represent 50 mm, 100 mm and 140 mm sample sizes respectively.
surface, while still having good oxygen supply from the free surface to allow smouldering propagation.

The smoulder spread rates range from approximately 5 mm min\(^{-1}\) to 10 mm min\(^{-1}\) for smouldering at heat fluxes just above the critical heat flux and increase up to around 25 mm min\(^{-1}\) before the onset of flaming. The error in the spread rate is estimated to be ±25% due to the uncertainty of ±2.5 mm in the thermocouple placement in the foam. The spread rates measured for smouldering are in agreement with the work of Anderson et al. [5] using a similar experimental set-up but are higher than those generally found for unassisted smouldering [3, 17]. The higher spread rate could be explained by the energy assistance provided by the heater.

Figure 12 shows the spread rate as a function of depth for smouldering in a 100 mm sample at heat fluxes of 8 kW m\(^{-2}\), 26 kW m\(^{-2}\) and 33 kW m\(^{-2}\). There is a steady increase in smoulder spread rate as the heat flux is increased. The spread rates between 15 mm and 25 mm depth are 6 mm min\(^{-1}\), 18 mm min\(^{-1}\) and 22 mm min\(^{-1}\) respectively. A strong dependence on depth is seen with spread rates decreasing to 3 mm min\(^{-1}\), 8 mm min\(^{-1}\) and 18 mm min\(^{-1}\) at a depth of 50 mm. The spread at 16 kW m\(^{-2}\) and 33 kW m\(^{-2}\) reaches 95 mm; however, the spread rate has reduced to 4 and 13 mm min\(^{-1}\), respectively. The same trend is seen for the 50 mm and 140 mm samples; however, for the 50 mm samples, the maximum spread depth is 75 mm.

It is known that the ignition of smouldering combustion is controlled by heat transfer and kinetics and the spread is controlled by the availability of oxygen and heat losses [6, 8]. Therefore, this decrease could be explained by the decrease in available oxygen as the smoulder propagates deeper into the sample. This means
that oxygen has to diffuse further before it reaches the reaction front and limits the spread rate. This effect has been previously reported by Palmer [22] and Krause and Schmidt [11]. Near the free surface, there is a large supply of oxygen and the reaction is dominated by the heat supplied by the heater.

4. Discussion

4.1. Sample Size

The sample size has been shown to have a strong impact on the smouldering and flaming ignition of polyurethane foams. The critical heat flux for smouldering and flaming are shown to decrease as the sample size is increased. This relationship appears to become less strong for larger samples. The behavior of the 50 mm samples is seen to differ significantly from that of the 100 mm and 140 mm samples. This could be due either to the small surface area available for oxygen diffusion to the sample which would limit the reaction rate, or increased heat losses to the surroundings for the 50 mm sample which has a higher surface area to volume ratio than the larger sizes. Unfortunately, it is not possible to conclude further using the data obtained from the current experimental set-up.

4.2. Transition to Flaming

The transition between smouldering and flaming ignition is a complex phenomenon marked by rapid changes in spread rate and peak temperatures. There are two possible mechanisms for this transition: gas phase ignition or the transition
from smouldering to flaming. The gas phase ignition may be due either to auto-
ignition or piloted ignition at the cone heater which, at high heat fluxes, operates
at temperatures exceeding 700°C.

Transition from sustained smoulder to flaming was not observed in any of the
experiments. However, it was not possible to conclude on the possible transition
from incipient smouldering to flaming.

There are two reasons that the transition from incipient smouldering to flaming
would not be observed with the current diagnostics set-up: flaming ignition would
take place at a much faster time scale than that required for a smouldering signa-
ture to develop in the temperature data, and transition would occur on a thin
layer within the foam, which the spatial temperature resolution used here (10 mm)
would be too coarse to capture.

Bustamante et al. [20] found the minimum heat flux for flaming auto-ignition of
the same PU foam as used in this work was 35 kW m⁻² (compared to 9 kW m⁻²
for piloted ignition). These experiments were carried out in the cone calorimeter
using samples with square cross section of 100 × 100 mm and height of 50 mm.
This heat flux for auto-ignition agrees well with the range found here for flaming
ignition for the same sample size (32 kW m⁻² to 37 kW m⁻²). This suggests that
the mechanisms leading to flaming ignition in [20] are the same as the mechanisms
in this work. Bustamante et al. concluded that flaming ignition was an auto-igni-
tion process but did not consider smouldering and transition to flaming as a pos-
sible mechanism, or the hot surfaces of the cone acting as a pilot.

Putzeys et al. [17] found that a minimum heat flux between 8 kW m⁻² and
8.75 kW m⁻² will result in the transition from smouldering to flaming of a
50 mm × 50 mm × 125 mm sample of PU foam with internal forced flow and
sample sides heated to 200°C by external means. This value is much lower than
that measured here for 50 mm size (45 kW m⁻² to 48 kW m⁻²), but the heating of
the samples in [7] sides and the increased air flow dramatically enhance the smoul-
dering combustion, increasing the likelihood of transition to flaming.

Using the temperature data collected and comparing the tests here to those
available in the literature, it is not possible to provide a definite mechanism for
flaming ignition. Further investigations using enhanced apparatus such as higher
thermocouple resolution to capture the smoulder reaction in more detail, gas anal-
ysis, high speed cameras, advanced flame detection techniques or changing the
separation between the sample and the cone heater but keeping the heat flux con-
stant could assist in the determination of the process leading to flaming ignition.

5. Conclusions

The effect of sample size on radiant smouldering ignition had been theoretically
proposed before but this is the first time that it is demonstrated experimentally.
The onset of smouldering ignition was observed in the ranges 13–14 kW m²,
8–9 kW m², 7–8 kW m² for 50 mm, 100 mm and 140 mm sample sizes respec-
tively. The onset of flaming ignition was observed in the ranges 45–48 kW m²,
32–37 kW m², 30–31 kW m², respectively.
Smouldering spread rates are measured in the range from 3 mm min\(^{-1}\) to 25 mm min\(^{-1}\) and found to be a strong function of the heat flux and depth of the smoulder front from the free surface. Due to significant variability in the measurements, there is no clear effect of the effect of sample size on the spread rate. However, it is observed that the 50 mm samples consistently show lower spread rates for a given heat flux than the 100 mm or 140 mm samples.

The onset of smouldering ignition is observed at significantly lower heat fluxes than flaming (approximately 70–80% lower). This offers a route to initiate flaming sources by the mechanism of transition from smouldering to flaming. Critical heat fluxes for both smouldering and flaming ignition increase with the sample size, with smouldering ignition being significantly more sensitive to sample size than flaming ignition under the range studied. Since most standard testing is conducted a one single sample size on the order of 100 mm, the observation reported here that the lowest flammability is observed for the largest samples could have implications for testing procedures and translation of small-scale testing to real applications in the built environment.

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