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Methodology for Estimating Pyrolysis Rates of Charring Insulation Materials using Experimental Temperature Measurements

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Abstract

This paper presents the application of a simplified method to estimate pyrolysis rates from rigid closed-cell cellular plastics by means of experimental temperature measurements. These materials are extremely effective in meeting energy efficiency goals in buildings and their safe use should also be enabled and optimised by undertaking comprehensive fire safety analyses. The proposed methodology consists of determining the mass loss as a function of the thermal evolution by applying a mass conversion directly using thermogravimetric data under non-oxidative conditions. In order to verify this simplified method, an experimental programme based on 100 mm thick samples of rigid polyisocyanurate foam was conducted using a Cone Calorimeter, obtaining measurements of mass loss and temperature within the core of the material. A Monel plate was used on top of the sample in order to represent a simpler boundary condition by eliminating the smouldering process of the charred material. Although the pyrolysis rates using this methodology did not provide a perfect fit with experimental data, they showed similar trends, with a slightly delayed prediction but still accurate magnitude. This methodology presents potential for fire safety engineering applications in two domains: (1) as a complementary technique to improve the interpretation of results from standard and ad-hoc testing, and (2) as a design technique for the evaluation of potential heat release contribution and gaseous emissions of assemblies incorporating insulation materials.

Keywords

Insulation materials; Fire Safety; Pyrolysis; Modelling; Performance-based design; Charring

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Nomenclature

\( k \) thermal conductivity (W·m\(^{-1}\)·K\(^{-1}\))
\( c \) specific heat capacity (J·kg\(^{-1}\)·K\(^{-1}\))
\( f \) non-dimensional fraction of remaining mass (-)
\( i \) number of element (-)
\( j \) number of time step (-)
\( k \) number of exposure area (-)
\( L \) thickness (m)
\( m \) mass (g)
\( \dot{m} \) mass flow (g·s\(^{-1}\))
\( \dot{m}'' \) mass loss rate per unit area (g·s\(^{-1}\)·m\(^{-2}\))
\( N \) maximum number of finite differences (-)
\( q \) heat (W)
\( t \) time (s)
\( S \) surface area (m\(^2\))
\( T \) temperature (K or °C)
\( x \) distance (m)
\( Y \) yield (g·g\(^{-1}\))

Greek letters

\( \alpha \) absorptivity/emissivity (-)
\( \Delta H_c \) effective heat of combustion (J·kg\(^{-1}\)·K\(^{-1}\))
\( \Delta t \) time step (s)
\( \Delta x \) finite difference thickness (m)
\( \rho \) density (kg·m\(^{-3}\))

Subscripts

\( 0 \) initial
\( cr \) critical
\( i \) of the difference i
\( net \) net/conductive
\( P \) pyrolysis
\( z \) species

Acronyms

HRR heat release rate
MLR mass loss rate
PIR rigid closed-cell polyisocyanurate foam
SIP structural insulated panel
1. Introduction

During recent decades sustainability has become one of the main drivers in building construction, resulting in highly thermally efficient buildings. Several techniques may be used to achieve the stringent energy efficiency requirements defined by the Energy Performance of Buildings Directive [1], e.g. thermal insulation within the building envelope, increased levels of air tightness, efficient heat recovery of the ventilation systems, reduction of thermal bridging and/or more efficient windows [2]. The intense use of thermal insulation is one of the primary targets due to the large surface area of the building envelope and the architectural aspirations. As a result, low thermal transmittances (U-values) are required, which can only be achieved by significantly increasing the thickness of insulation used.

Due to the multi-criteria nature of building design, stringent U-values clash with other desired design criteria such as efficient space usage and cost. Despite the large diversity of insulation materials in the market [3], under this competitive scenario closed-cell plastic foams have become an easy and cost-effective solution because of their relatively low thermal conductivity. The most common closed-cell insulation foams at present being used are rigid polyisocyanurate foams, commonly known as PIR, and phenolic foam. These materials are often provided as boards with a foil-facing on the surface and used for framing construction or masonry cavity walls; alternatively they can be embedded directly within linings, e.g. sandwich panels or structural insulated panels (SIPs) [4].

Despite the fact that these materials are extremely effective in meeting energy efficiency goals, their use should be also enabled and optimised by undertaking a comprehensive fire safety analysis, i.e. systems including insulation materials should be optimised while still ensuring life safety and property protection.

1.1. Fire performance of closed-cell plastic insulation materials

The fire performance of these materials has been studied by several authors at different scales [5–19]. Generally, these types of plastics are classified either as thermoplastics or thermosets. Thermoplastics (e.g. expanded polystyrene) exhibit melting behaviour, while thermosets (e.g. polyisocyanurate or phenolic foam) exhibit a charring behaviour, leaving a carbonaceous residue after pyrolysis. A complete description of the different mechanisms of thermal decomposition for these polymers is described by Witkowski et al. [5]. These mechanisms result in different fire performance, with a charring behaviour being more desirable due to the positive effect of the char layer on the reduction of the pyrolysis rate. Several authors have focussed their research at the material scale (e.g. thermogravimetry), looking at polymer formulations that
promote larger residue generation and endothermic reactions in the solid-phase [6–8]. These techniques of flame retardancy have been largely covered by Hull and Kandola [9]. However, the majority of research has focussed on the macroscopic material behaviour using bench-scale testing, thus concentrating on the ignition mechanism and release of heat from these materials [10–19]. More extensive experimental work covering different scales can be found in references Error! Reference source not found. and Error! Reference source not found.

Recently published work showed the relation between the thermal degradation at the material scale linked to the heat transfer phenomena within the solid material [20]. Rigid closed-cell polyisocyanurate and phenolic foam showed similar behaviour, i.e. materials that experience pyrolysis and char formation. The char layer reduces the heat transport to the pyrolysis front resulting in a slower propagation and lower pyrolysis rate. Typically, this insulating effect of the surface char layer limits the heating of virgin foam to several degrees per minute. Experimental results showed that this char is however highly vulnerable to surface oxidation (smouldering). The smouldering process was shown not to be self-sustaining due to the large heat losses under the specific experimental conditions. In addition, the closed-cell structure of the polymer restricted the air flow through the foam which was shown to be a key factor to limit self-sustaining smouldering [20]. In end-use conditions, the insulation materials are typically covered by a lining or a physical barrier, thus limiting the contact with the air, unless they are introduced in partial fill cavity walls. As a result, this smouldering behaviour is not expected under real fire conditions. Therefore, initially only pyrolysis should be considered as the primary hazardous event.

1.2. Fire safe design of insulation materials

It has been widely recognised that the organic polymer nature of closed-cell plastic foams may represent a fire risk in buildings [21–23]. The fire safe design of building assemblies including insulation materials has been classically based on a material classification and a pass-fail criteria frameworks, e.g. in the EU represented by the Euroclasses system [24] and the fire-resistance framework [25], respectively. The prescriptive nature of these frameworks however does not allow for a quantitative design to be carried out on the basis of the specific insulation fire hazards, and as a result it is not possible to quantify the associated fire risk [26].

Previous work demonstrated that the initiating hazard from this type of insulation material corresponds to the onset of pyrolysis [26]. After this is achieved, there is potential for the generation of a large amount of flammable gases that may be transported to the compartment fire, or alternatively may escape to areas away from the fire enclosure. The former may represent an increase in the heat release rate of the fire, while the latter may represent a life safety hazard for the occupants of the building due to the intrinsic toxicity of
the pyrolysis effluent. Current mitigation practices are thus based on the design of suitable thermal barriers that delay or cancel the onset of pyrolysis of the insulation material [27].

Whilst the previous approach stands out by its simplicity, which is easily achievable in engineering terms, it may be very conservative for some scenarios. A more accurate approach should rely on estimating the rates of pyrolysis from the insulation under specific fire scenarios. By determining the pyrolysis behaviour, the evolution of the hazard (potential contribution to the fire and generation of toxic species) can be quantified [28]. This approach requires a pyrolysis model which is able to accurately predict the thermal behaviour of the insulation. In the last decade, such pyrolysis models have been developed and validated [29–32]. These models tend to require a large number of parameters that are often unknown, necessitating inverse modelling techniques that can introduce significant compensation errors [33] and demand a great deal of expertise. An additional handicap on the use of these tools is that the thermal boundary condition under real fire scenarios is difficult to quantify. Consequently, if pyrolysis rates are to be quantified at an appropriate level for engineering design optimisation, simpler approaches are necessary.

1.3. Research aim

This work aims to assess the applicability of a simplified methodology for quantifying pyrolysis rates and temperature evolution from foil-lined closed-cell charring insulation materials under severe conditions of heat exposure. It focuses on small-scale experiments so as to reduce the uncertainty in the assessment, thus precisely controlling the thermal evolution and mass loss of samples under a heating regime that is close to one-dimensional. Despite the fact that characterising the pyrolysis represents a challenge due to the large number of material properties to be quantified, the presented simplified experimental approach still allows prediction of the hazard while keeping the method simple. The success of this approach will allow estimation of pyrolysis rates from this type of insulation in research-driven large-scale experiments and standard testing. Given that the pyrolysis rate represents the main physical variable determining the heat release contribution and yields of toxic species, the application of this methodology will help to improve current testing practices.

2. Experimental programme description

The experimental programme was designed to explore the applicability of simpler modelling approaches, and based on the use of the Cone Calorimeter apparatus [34] and thermogravimetric experiments. The Cone Calorimeter set-up was modified to remove the pilot spark and to enable heating of the sample by conduction from a metallic plate on the exposed surface. The main measurements consisted of mass loss and temperature within the samples, supported by visual observations. Four experiments were performed for each thermal exposure, two repetitions measuring only mass loss, and two repetitions taking
temperature measurements within the samples. The thermogravimetric data corresponded to those presented by the authors elsewhere [35].

2.1. Materials

The studied insulation material corresponded to a type of rigid polyisocyanurate foam previously described as PIRb elsewhere [20,35]. Samples with a surface area of 90 mm by 90 mm and 100 mm thick were tested for this series of experiments. The metallic plate as boundary element at the surface of the samples corresponded to a 6 mm Monel plate, painted with a high temperature optical black coating of known absorbivity ($\alpha = 0.92$, Medtherm Corporation®). The use of the plate presents a case study representative of a common end-use condition of insulation materials, as these are rarely installed uncovered but behind a lining. The oxidation rate is expected to be reduced or eliminated by using this methodology, therefore also reducing the complexity introduced by the smouldering process for future modelling purposes. A metallic plate (6 mm thick, Nickel 200) was used at the bottom of the sample as a heat sink. A sample with the protective foil layer removed and prepared for testing with the metallic plates on top and bottom is shown in Figure 1a.

![Figure 1a](image1.png)

![Figure 1b](image2.png)

Figure 1. (a) PIR sample prepared for testing. (b) PIR sample wrapped in aluminium foil and ceramic paper with metallic plates and thermocouples inserted into the centre of the sample through ceramic tubes. A special holder was designed to keep the thermocouple horizontal during the insertion.

2.2. Experimental set-up

The samples were wrapped with aluminium foil on the bottom and lateral sides, with the 6 mm Monel plate placed on the top and the 6mm Nickel 200 block at the bottom. Sample and plates were wrapped in two 3 mm thick layers of ceramic insulation paper. The purpose of the aluminium foil was to prevent air penetration into the sample from the sides, as the ceramic paper is a porous element, to promote one dimensional pyrolysis. The real set-up and a schematic drawing of this are shown in Figure 1b and Figure 2.
In order to provide a well-characterised experimental set-up to allow modelling to be undertaken, the characterisation of the boundary condition at the back face of the material was achieved by using the 6 mm Nickel 200 plate at the bottom of the samples. This approach was described by Carvel et al. [36], who recommended the use of a heat sink for material characterisation purposes. Using a metallic plate on top would act as a dummy surface temperature sensor, although contact resistance effects may induce a thermal gradient between sample and metallic plate. This is discussed in subsequent sections.

Several levels of irradiation from the radiant heater were used (25, 45 and 65 kW·m$^{-2}$). The heat fluxes were selected such that different rates of pyrolysis would be achieved. Measurements of temperature were taken within the sample by using 1 mm bead N-type thermocouples. The temperature of the metallic plates on the top and bottom was also measured, but with 1.5 mm bead K-type thermocouples. Thermocouples were installed at various depths at the centre of the section (1, 10, 20, 30, 40, 60, 80 mm) parallel to the exposed surface to reduce the error in the thermocouple reading, which is recommended for materials of particularly low conductivity [37,38]. Ceramic tubes were used to insert the thermocouple into the sample, so as to secure the location of the thermocouple for multiple experiments. Additionally, the exact location of the thermocouples could be visually identified after testing. No temperature correction was considered by the heat losses introduced by the thermocouple. The positioning of the thermocouples is shown in Figure 2. A summary of the conditions for all the performed experiments is presented in Table 1.

Figure 2. Schematics of sample preparation.
Table 1. Summary of performed experiments

<table>
<thead>
<tr>
<th>Material characteristics</th>
<th>Configuration</th>
<th>Incident radiant heat flux range /kW·m⁻²</th>
<th>Measured parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIRb</td>
<td>Nominal sample size: 90mm x 90mm x 100mm</td>
<td>25, 45, 65</td>
<td>(1) Mass loss (2 repetitions)</td>
</tr>
<tr>
<td></td>
<td>Top boundary condition: Monel 400 plate (6mm)</td>
<td></td>
<td>(2) In-depth temperature (2 repetitions)</td>
</tr>
<tr>
<td></td>
<td>Wrapping: 2 layers of ceramic paper + 1 layer of aluminium foil</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Back boundary condition: Nickel 200 plate (6mm) + Ceramic board (25mm)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orientation: Horizontal</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pilot: No pilot igniter</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Manufacturer-claimed density: 32 kg·m⁻³</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Average measured density: 33.0 ± 0.71 kg·m⁻³</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Estimated thermal inertia: 6.5·10⁻⁴ W²·s·K⁻²·m⁻⁴ [27]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

3. Experimental results and discussion

Figure 3a shows the normalised mass for PIR samples with a Monel plate on top under three constant levels of irradiation (25, 45 and 65 kW·m⁻²). The normalised mass is obtained by dividing the mass of the sample at any time by the initial mass \(m_0\). Vertical dashed lines indicate the time at which the effluent of pyrolysis gases through the edge of the sample auto-ignited. Results from duplicates show good agreement, with the major discrepancies observed for the highest heat flux after ignition is observed. This inconsistency in the results is however expected due to the behaviour of the pyrolysis effluent for each experiment, resulting in extra heating of the surface metallic plate when ignited. The sample residue obtained after 1800 s of a heat exposure of 25, 45 and 65 kW·m⁻² is approximately 88%, 75% and 60%, respectively.

Figure 3b shows the mass loss rate (MLR) per unit area corresponding to the mass loss presented in Figure 3a. The shape obtained for the three heat fluxes is qualitatively similar, with a MLR peak followed by a decay, as characteristic of charring materials [39]. The MLR peak increases in magnitude and shifts to lower times with increasing heat fluxes. Peaks of MLR for 25, 45 and 65 kW·m⁻² are approximately in the range 0.4-0.5, 1.0-1.2 and 2.0-2.4 g·m⁻²·s⁻¹, respectively. Considering a heat of combustion for the pyrolysis gases of 13.22 kJ·g⁻¹ as presented by Hidalgo [35], the heat release rate per unit area of these peaks correspond to 6.6, 15.9 and 31.7 kW·m⁻², which are fairly moderate values. Therefore, a significant contribution to the heat release in a compartment fire from the insulation is only to be expected if a large surface area is exposed and this is limited to the early stages.
Figure 3. (a) Normalised mass for experiments at 25, 45 and 65 kW·m\(^{-2}\). Vertical dashed lines indicate the auto-ignition of the pyrolysis effluent. (b) Mass loss rate per unit area. The sample residues for experiments shown in Figure 3 are presented in Figure 4, with a cut through the centre-section. The sample sections show three different regions of discoloration corresponding to the char after pyrolysis (carbonaceous region), reaction zone where pyrolysis process is ongoing (orange region), and virgin material (light yellow/grey region). The regions of discoloration are fairly uniform along the width of the sample, indicating a heating regime similar to 1D. This uniformity is less clear in Figure 4a, corresponding to the sample tested at 25 kW·m\(^{-2}\). These residues are coupled with the temperature profile during the quasi-steady\(^3\) state after 1800 s, obtained from experiments using thermocouples. It should be

\(^3\) The term “quasi-steady state” used throughout this paper refers to the stage in which the rate of temperature increase within some sections of the sample is sufficiently low that the net heat flux at the sample surface has achieved an asymptotic behaviour, close to a constant heat flux that defines a steady conduction.
noted that the temperature measurements are presented as the original locations in the sample before testing. The samples tested at 25, 45 and 65 kW$\cdot$m$^{-2}$ presented a shrinkage after 1800 s of about 0, 5 and 10 mm, respectively. For ease in the visualisation, the image of the residue has been adapted to the original sample size of 100 mm to fit into the same scale of the temperature-depth diagrams.
Figure 4. PIR residue after 1800 s of heat exposure of (a) 25, (b) 45 and (c) 65 kW·m$^{-2}$. The maximum temperature achieved for each depth is presented on top of each cut presented as background.
Figure 5 shows the temperature measurements obtained for experiments with thermocouples under 25, 45 and 65 kW·m$^{-2}$. Duplicated experiments are shown as dashed lines. In general, good repeatability is obtained for experiments at 25 and 45 kW·m$^{-2}$, while for 65 kW·m$^{-2}$ this is not that good for thermocouples in the 10 mm and 20 mm in-depth position during the transient state, and also for the top plate. The temperature at the top plate experiences a sudden increase due to the auto-ignition of the pyrolysis effluent. Then, the temperature decreases significantly, with a noisy reading, thus indicating a bad contact between thermocouple and plate, which was later corrected.

The readings presented in Figure 5 indicate a clear effect of the contact resistance during the experiments. During the transient state the temperature difference between plate and first position within the sample (1 mm) is clearly noticeable with a difference of up to 200 °C. However, once the quasi-steady state is reached, the difference drops to only 20 °C. This is a reasonable result, as the net heat flux through the surface of the sample is large during the transient state, reducing continuously until the quasi-steady state is reached. Given a constant thermal resistance of the contact, a larger heat flux will result in larger temperature gradients. While these considerations about the contact resistance are important for heat transfer modelling, for the present analysis they have few implications as a reading near the surface is also available.

Additionally, the obtained measurements indicate that the smouldering process has been successfully mitigated by using the metallic plate and aluminium foil, as the temperature evolution follows the trend of an apparently inert solid under a constant irradiation level and heat losses. The critical temperature proposed by Hidalgo et al. [27] for this material (PIR) is 300 °C, which fundamentally represents the onset of hazard (pyrolysis) and corresponds to the primary failure criterion to be considered for the fire safe design of assemblies including insulation. This value is plotted as a horizontal line, showing that samples exposed to 25, 45 and 65 kW·m$^{-2}$ achieve the critical temperature at the 1 mm in-depth thermocouple at about 520, 285 and 220 s, respectively. These times slightly correspond to the period prior to the maximum increase in MLR before the peak, thus validating the conservative definition of critical temperature for charring materials proposed by Hidalgo et al. [27].

Figure 6 (solid lines) shows the propagation of the front at 300 °C (assumed to correspond to the pyrolysis front) obtained for the three cases presented in Figure 5 by interpolating the temperature profile for each position. Due to the imprecise positioning of some thermocouples and/or density of temperature measurements within the insulation core, it is observed that the position versus time presents a change of curvature, which otherwise would not be expected e.g. at 500 s, 20 mm depth during the 45 kW·m$^{-2}$ heat exposure. Complementary to this, the first derivative of this function that represents the spread rate of the front at 300°C is plotted as a dashed line. A maximum spread rate of 2, 5 and 6 mm·min$^{-1}$ is observed for
25, 45 and 65 kW·m⁻², respectively. Consistent with the data of normalised mass, the spread of the pyrolysis front experiences an attenuation/decay due to the charring nature of the foam, which is fundamentally a consequence of a reducing net heat flux at the pyrolysis front. The fact that the char layer is protected by the metallic plate, and therefore not consumed by oxidation, allows the net heat flux at the pyrolysis front to keep decreasing as this progresses in-depth.
Figure 5. Temperature profiles for a heat exposure of (a) 25, (b) 45 and (c) 65 W·m⁻². Repetition shown as dashed lines.
Figure 6. Position of the front at 300 °C (solid lines) and spread rate of this front (dashed lines) for experiments at 25, 45 and 65 kW-m². Interpolated values computed based on data presented in Figure 5.

4. Modelling analysis

4.1. Principles for the simplified pyrolysis rate assessment

The simplified pyrolysis model is based on the approach already proposed by Hidalgo et al. [40]. This approach consists of a two-step decoupled analysis, first solving the heat transfer problem and then followed by the estimation of remaining mass and pyrolysis rates. Despite the fact that effective thermal properties could be obtained to characterise the PIR thermal evolution, in this work modifications are made and the first step is bypassed experimentally without having to solve the diffusion heat transfer within the solid-phase. The approach then consists in determining the mass loss as a function of the thermal evolution. The sample is considered as the space domain \( x = L \) (m) divided into \( N \) finite differences of thickness \( \Delta x_i \) (m), with \( i \) each of the finite differences. As for the analysis in the previous section, the temperature evolution for each finite difference is obtained by linear interpolation. Given that the method is also discretised in time, each time step is defined as \( j \) and considered as \( \Delta t \) (s). Then, the normalised sample mass for the time step \( j \) is obtained as the following expression representing an integration over the space domain:

\[
\bar{m}_j = \frac{\sum_{i=1}^{N}(\bar{m}_i \cdot \Delta x_i)}{L} \tag{1}
\]

where \( \bar{m}_i \) is the normalised mass of the finite difference \( i \), which is approximated directly as a function of the temperature \( f(T) \):
\[ \bar{m}_i = f(T) \]  

The function \( f(T) \) establishes the fraction of remaining mass as a function of the temperature in that finite difference, varying from 0 to 1. To simplify this function and remove uncertainty associated with fitting of Arrhenius parameters, which depend on the temperature and the concentration/diffusion of oxygen [41,42], \( f(T) \) is defined by direct reference to TGA results under sufficiently low heating rates. The TGA curves presented in Figure 7 correspond to PIR from the same manufacturer obtained in a non-oxidative atmosphere and heating rates of 2.5 and 20 °C·min⁻¹ [4]. The normalised mass loss rate can be obtained by deriving the mass loss over time, which in a discretised form corresponds to the increment of the normalised mass between time steps divided by the time step. The mass loss rate per unit area can then be calculated by considering the density of the virgin material \( \rho_0 \) (kg·m⁻³) and the thickness of the sample \( L \) (m):

\[ \bar{m}_P^{ij} = \rho_0 \cdot L \cdot \frac{\bar{m}^{i-1} - \bar{m}^j}{\Delta t} \]  

where \( \bar{m}_P^{ij} \) is the mass loss rate per unit area (kg·m⁻²·s⁻¹), or equivalently the rate of pyrolysis per unit area because no oxidation is considered.

![Figure 7. Normalised mass of PIR from the same manufacturer obtained by thermogravimetry under a nitrogen atmosphere at 2.5 and 20°C·min⁻¹ [4].](image)

4.2. Results

Figure 8 shows the experimental and modelled normalised mass loss rate considering the temperature profiles presented in Figure 5. Normalised values of mass loss rate are compared so that the error due to inaccuracy in the dimensions, and therefore density calculation for the modelled MLR, can be minimised. The experimental normalised MLR is obtained by differentiating the normalised mass curves with respect to time (as shown in Figure 3a), while the modelled MLR is obtained by integrating the \( f(T) \) function for
the whole thickness for each time step as shown in Eq. 1 and 2, and then differentiating with respect to time.

It is observed that the model does not produce a perfect fit with the experimental data. This is however not surprising due to the deliberate simplicity of the proposed approach. Nevertheless, it is clear that the primary mechanisms that characterise the pyrolysis from PIR under different heating scenarios are fairly represented, since the results follow similar trends. The most obvious inaccuracy presented by the model is the delay between experimental and model results. This could be partially attributed to a bad adjustment of the time lines for experiments with and without thermocouples. Other factors that could contribute to this delay are an insufficiently high density of thermocouples near the surface of the material and/or the linear interpolation method. The transport time of the pyrolysis gases could also be a factor, although the time scale is expected to be much lower, of the order of seconds. In any case this is a drawback that can be easily corrected and/or calibrated without affecting the outcomes of the assessment.

Another clear conclusion from the results presented in Figure 8 is that the TGA curve for which the model better predicts the experimental results is the one with the highest heating rate used, i.e. 20 °C·min⁻¹. Figure 9 shows the heating rate experienced for various locations (1 and 4 mm) at 25 and 65 kW·m⁻². The derived maximum heating rate is around 140 °C·min⁻¹, while for other regions and heat fluxes the heating rate does not go over 60 °C·min⁻¹. The slight overestimation of the modelled MLR is consistent with these results, as 20 °C·min⁻¹ is not as high as the heating experienced at certain locations. However, the observed heating is obviously not constant, with expected average values closer to the 20 °C·min⁻¹ threshold. Despite the fact that using this heating rate as input for the function $f(T)$ may lead to a slight overestimation of results, in engineering practice this could still be a conservative and practical approach. As a matter of fact, the results using a lower heating rate do not show significantly large overestimations.

Figure 8c shows that the model based on the data from one of the repetitions with thermocouples at 65 kW·m⁻² presents a clear and more significant overestimation of the MLR. This is probably mainly due to inaccurate positioning of the thermocouples for this particular experiment, which did not use a stiffening system as presented in Figure 1b. This highlights the importance of the position and density of temperature measurements in practice.
Figure 8: Experimental and modelling results for (a) 25, (b) 45, and (c) 65 kW·m$^{-2}$. Experimental curve corresponds to the average between two repetitions, and the shading is the maximum deviation from repetitions to the average.
Figure 9. First derivative of temperature measurements at 1 and 4 mm at 25 and 65 kW·m\textsuperscript{-2}.

4.3. Limitations and model uncertainties

Oxidative conditions

The proposed model is based on the assumption that no smouldering occurs. Therefore, the applicability of this approach is only valid for non-permeable materials that do not allow oxygen transfer through the matrix, thus the thermal decomposition corresponds to non-oxidative conditions as presented in Figure 7. If oxidative conditions are produced on the top, exposed surface, the calculated rate of pyrolysis is expected to still be reasonably reproduced, however the total mass loss rate would be under-predicted as the char oxidation would be neglected.

The shrinking behaviour of the material would lead to the creation of a gap between lining and insulation. Once this is achieved, air flow may be expected within the gap. After this has occurred the smouldering process may become a relevant hazardous event that would increase the heat flow through the insulation reaction zone.

Heating rate

While it is normally admitted that the heating experienced by burning fuels is much larger than heating rates obtained by thermogravimetry [43], this is clearly not the case for charring materials such as PIR. The effectiveness of the approach then resides in the low and reducing heating rates experienced by the reaction and virgin zones, as shown in Figure 9. The results presented in the previous section indicate that a heating rate of 20 °C·min\textsuperscript{-1} provides a reasonable accuracy for all the heating scenarios studied and the particular thickness of 100 mm. The trend in Figure 8 indicates that increased heating rates could provide better results, since 20 °C·min\textsuperscript{-1} is a heating rate lower than those observed in Figure 9 during the peaks of MLR; however the improvement appears to be not very substantial.
Thermal interpolation

As shown in Figure 8c, an accurate position of the thermocouples and their spacing is essential to obtaining sensible results. For the present case of 100 mm thick samples, 10 mm spacing for regions near the surface and 20 mm for regions far from the surface are able to provide sufficiently good results. While smaller spacing may result in greater accuracy, a system to secure the thermocouple as presented in this experimental programme seems essential, as the error in spacing can easily be of order ± 3 mm when thermocouples are inserted without a stiffening system, due to the friable nature of the foam. In addition, the thermal mass of an increased number of thermocouples may result in a 'heat sink' effect, leading to premature quenching of the reaction.

The interpolation method used to obtain thermal evolution between thermocouple measurements was linear. The accuracy of this approach is proportional to the density of thermocouples. Despite the simplicity of the approach, it seems to provide accurate results.

Heat transfer dimensionality

The accuracy of the results from this experimental programme also depend on the one-dimensionality of the heat transfer. Observations of the colour change of the experimental sample residues qualitatively indicates that the 1D assumption seems to be fairly correct. While this could be an issue for the presented experimental programme, since the wrapping material had higher conductivity than the samples, in real scenarios however this assumption is rather controlled by the uniformity of the heating boundary condition. For that case, the accuracy would then be limited by the density of measuring points over the surface area of the building assembly being studied.

5. Applicability for fire safety engineering

Since the primary fire hazard from these types of insulation can be associated to the pyrolysis process, the main parameter to be quantified is the rate of pyrolysis gas release. Despite the fact that there is large uncertainty with regard to the location, conditions and instant at which these will ignite, due to, for instance, the ventilation conditions of the construction system, the conservative approach is to assume that these would instantaneously contribute to the fire. This way the risk can be quantified more easily.

The presented method, although not extremely accurate, presents a reasonable level of precision for engineering purposes where the degree of uncertainty in other parameters is already high. Two clear applications can be found for the present model as: (1) pyrolysis estimation for fire testing such as large-scale experiments or standard testing, and (2) for quantitative design purposes.
5.1. Model testing

The presented simplified method can be used to develop a model of the pyrolysis behaviour under well-defined testing conditions. The concept consists of running ad-hoc and/or standard fire testing including a series of thermocouples to allow an *a posteriori* quantification of the pyrolysis behaviour from the insulation. This approach would also allow quantification of the effectiveness of various protection systems, to allow an optimised design solution to be generated. The provision of pyrolysis predictions can complement heat release rate calculations and measurements of gas species from the generated smoke. Within this scope, error bars need to be acknowledged, based on the model limitations and testing conditions noted previously. Clearly, if consistent data are used, expected errors can be quantified and delimited. The potential of this approach resides in the low-cost solution for improved product development, thus reducing costly research based on full-scale testing.

5.2. Hazard quantification

The proposed methodology can lead to evaluation of designs, by which the fire hazard from insulation materials can be quantified explicitly if a series of assumptions and hypotheses are established. A diagram describing the application of the methodology for design purposes is presented in Figure 10. The approach consists of evaluating the time to achieve a potential hazardous heat release contribution and gaseous emissions from the insulation.

The first step is based on the definition of effective thermal properties and initial thickness for lining and insulation. Then, a series of fire scenarios defined as the conditions of heat exposure (thermal boundary conditions), their respective area of exposure, and exposure time have to be proposed. Next, the thermal evolution of the system lining-insulation has to be estimated for each boundary condition by using a heat transfer solver. For simplicity, the problem can be simplified as a one-dimensional problem and a perfect contact can be assumed between insulation and lining, which are conservative assumptions. At this stage practitioners can either apply the simplified methodology based on a critical temperature proposed elsewhere [26], or alternatively apply the uncoupled pyrolysis model presented in previous sections in order to estimate the pyrolysis rate for each area of exposure. If the former is applied, the failure time of the insulation system is defined as the time when the insulation reaches the critical temperature at the surface. If the latter is applied, a total rate of pyrolysis gas generation needs to be calculated as the sum from each of the exposure areas. The potential heat release contribution can be obtained by multiplying the generation rate by the corresponding effective heat of combustion, while analogously the potential gaseous emissions can be obtained by multiplying by the corresponding yields. The failure time then can be defined as the time to reach a critical value of HRR or emission concentration.
While the calculation of gaseous emissions represents an ambitious task, as these strongly depend on conditions such as oxygen concentration and temperature [44], a series of hypotheses can be set if further toxicity assessments are pursued. For instance, these values can be used as inputs for CFD modelling in order to estimate fractional effective concentrations/doses for tenability assessments [45]. Similarly, potential HRR contributions can be used for tenability assessments in fire (zone/CFD) models.

The potential of this approach resides in the fact that the data required to develop these quantifications can be obtained by using bench-scale tests (for instance, thermogravimetric data or material properties such as thermal properties, yields or heat of combustion) that are often readily available from manufacturers, or by using values presented in the literature.
Figure 10. Evaluation process to determine the time to reach unsafe conditions for a specific wall typology and fire scenario (boundary condition).
6. Conclusions and future work

This paper has presented the application of a simplified methodology to estimate pyrolysis rates from charring insulation materials such as rigid polyisocyanurate foam based on experimental temperature measurements. In order to verify the proposed method, an experimental programme consisting of 100 mm thick samples of rigid polyisocyanurate foam was conducted using a Cone Calorimeter, obtaining measurements of mass loss and temperature within the core of the material. A Monel plate was used on top of the sample in order to represent a simpler boundary condition by eliminating the smouldering process of the charred material.

The proposed approach, although not showing a perfect fit with experimental data, has been proved to provide a reasonably good prediction of the pyrolysis rate from a rigid closed-cell polyisocyanurate foam under different scenarios of heat exposure using constant levels of irradiation (25, 45 and 65 kW·m⁻²), and a sample thickness of 100 mm. The simplified methodology is based on the direct use of TGA data to obtain mass loss measurements relying on temperature readings. Since this methodology is limited to non-oxidative conditions, the method must be applied for configurations where the insulation is covered by a lining or barrier. The use of TGA data with a heating rate of 20 °C·min⁻¹ and under a non-oxidative atmosphere has been shown to provide a good accuracy, with a slight overestimation in the modelled normalised mass loss rate. The position and density of temperature measurements has been highlighted to be one of the most important factors for achieving accurate results.

The presented method presents a reasonable level of precision for engineering purposes where the degree of uncertainty in other parameters is already high. This approach presents potential for fire safety engineering applications in two domains: (1) as a complementary technique to improve the interpretation of results from standard and ad-hoc testing, and (2) as a design technique for the evaluation of potential heat release contribution and gaseous emissions of assemblies incorporating insulation materials. The former approach can be used to allow a better characterisation of the fire performance of the insulation, without the necessity of developing complex numerical models. The latter approach can be used to evaluate designs by which the fire hazard from insulation materials can be quantified explicitly.

Since the approach presented in this paper is limited to experimental measurements of temperature, it may be insufficient for general design purposes, as it is scenario dependent. The main potential of the proposed methodology is that it would allow faster and more effective product development, thus reducing costly research based on full-scale testing. Therefore, future work should focus on determining effective thermal properties that could reproduce the thermal behaviour of the material and treat the problem as a
two-step uncoupled analysis, as shown elsewhere [40]. The experimental work presented within this paper can further be used for the determination of these effective model parameters.

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