# Infrared thermography as an operando tool for the analysis of catalytic processes: how to use it?

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# **Abstract**

Infrared (IR) thermography is a powerful tool to measure temperature with high space and time resolution. A particular interesting application of this technology is in the field of catalysis, where the technique can provide new insights into dynamic surface reactions. This paper presents guidelines for the development of a reactor cell that can aid in the efficient exploitation of infrared thermography for the investigation of catalytic and other surface reactions. Firstly, the necessary properties of the catalytic reactor are described. In fact, special equipment must be developed to ensure the realization of true *operando* conditions in the IR thermography experiments. Here, we provide the guidelines to assemble a chemical reactor with an IR transmitting window through which the reaction can be studied with the infrared camera. Secondly, we analyze the requirements towards the catalytic system to be directly observable by IR thermography. This includes the need for a catalyst that provides a sufficiently high heat production (or absorption) rate. We present selected examples of catalytic reactions that can be monitored by IR thermography, showing the potential of the technology in revealing transient and steady state chemical phenomena.

**Keywords**: IR thermography, operando methods, heterogeneous catalysis, exothermic reactions, transient phenomena

# 1. Introduction

Over the last decades, *operando* spectroscopy -i.e. spectroscopy coupled simultaneously with measurements of catalytic activity- significantly improved the knowledge of catalytic systems in industrially relevant conditions [1]. *Operando* spectroscopy is aimed at the determination of the variation of a wide range of catalyst properties during chemical reactions, including catalyst structure, oxidation state and reactivity. A special example of *operando* spectroscopic tool is infrared (IR) thermography. As a measurement technique, IR thermography operating principle is based on the detection of the IR radiation emitted by every object with a temperature above 0 K [2]. The measured radiation can be directly correlated with the temperature of the object, allowing for a precise and fast determination of the temperature [3]. The methodology is widely employed outside the field of catalysis, for example for the determination of temperature in medical applications or for

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the analysis of heat losses in buildings [4-6]. The main advantages of IR thermography are the relatively high resolution (each pixel is fundamentally one temperature sensor) and high measurement rate which modern IR cameras offer. Commonly, an IR camera can provide several thousands of data points in its field view, with a measurement frequency of dozens of Hz [7]. In particular, cooled IR cameras can achieve a very high sampling rate (above 100 Hz), while uncooled cameras, although cheaper, can record images with lower frequency [8]. Therefore, the main advantage of IR thermography over other temperature measurements, such as thermocouples and pyrometers, is the high spatial resolution and contactless investigation of reactions [9,10]. Furthermore, the temperature measurement with devices placed inside the reactor (i.e. with thermocouples) inevitably interferes with the experiments [11]. Despite these evident advantages, the examples in literature of systems for the systematic use of IR thermography as an operando tool for the study of chemical reactions are scarce. The main limitation to the development of the technique is related to the need of apposite reactors for this scope. In particular, the installation of an appropriate IR transmitting window in a chemical reactor may be challenging. Significant examples of application of IR thermography as operando method include the study of exothermic reactions, the determination of the temperature rise in structured catalysts and the utilization in high-throughput experimentation [9,12–17].

In this paper, we propose a systematic approach to apply IR thermography in operando studies. This kind of systematic experimental layout is currently lacking in literature and therefore we mainly aim at filling this gap in the scientific literature. We offer an overview of the main advantages and limitations of the methodology, showing how the selection of the reaction to investigate and the design of an appropriate experimental setup are interconnected. First, we define the requirements of the IR thermography system and elucidate the main possibilities to manufacture a reaction cell that allow the observation of the reaction without influence of parasitic phenomena. Furthermore, we discuss the selection of a suitable IR transmitting window, which can depend on the camera system selected (detection wavelengths) and the expected reaction conditions. Secondly, we define appropriate methods to systematically extract the significant information from the IR thermography experiments. In particular, we underline some general rules for the selection of the catalytic system, with respect to heat emissions and the reaction phenomena expected. Additionally, we derive some additional considerations about how to handle the experiments, to avoid contamination of the results by side effects (e.g. by mismanagement of the external heating). This involves the selection of appropriate additional analytical tools to correctly integrate the thermal images with concentration profiles and further information concerning the catalyst. Finally, we expose selected experimental case studies to validate the application of the methodology and to elucidate the additional information IR thermography can bring to operando studies. These examples involve the catalytic CO<sub>2</sub> hydrogenation to methane, which proved to be an ideal case study for the development of IR thermography as operando tool for the investigation of heterogeneously catalyzed chemical reactions. In fact, the CO<sub>2</sub> methanation, or Sabatier reaction, is a highly exothermal reaction following the stoichiometry:

$$CO_2 + 4H_2 \leftrightarrow CH_4 + 2H_2O$$
  $\Delta H_0^R = -164.9 \, kJ/mol$  (1)

The reaction is enabled and kinetically enhanced by different metallic catalysts [18]. Literature reports  $Ru/Al_2O_3$  at temperatures between 200-400 °C as one of the highest performing catalysts in the Sabatier reaction [19,20]. During the activation and steady state operation of the Sabatier

reaction, highly localized reaction effects can occur. Such effects include, for example, rapidly moving reaction hotspots during the reaction activation or heat gradients due to gas concentration and reaction rate differences during steady state operation. IR thermography proved to be a potent method to investigate these phenomena, which would be difficult to identify with the standard temperature measurement devices [21]. The results here elucidated can be compared with several other studies available in literature, which applied IR spectroscopy in the analysis of the Sabatier reaction [9,15,17,22–25].

# 2. Design of the cell for operando IR thermography

#### 2.1. Selection of the IR camera

IR thermography cameras are commercially available and typically record in two different IR spectra ranges: 3  $\mu m$  to 5  $\mu m$  for cooled detectors such as Indium antimonide (InSb) and 7  $\mu m$  to 17  $\mu m$  for uncooled detectors such as Microbolometer Focal Plane Arrays [26,27]. According to the black body spectral emittance diagram, reported in figure 1, we can correlate the temperature of a body with the wavelength of the emitted radiation and calculate the intensity of the IR radiation. Therefore, in catalysis applications, the characteristics of the experimental apparatus must be chosen according to the properties of the system to investigate. A decision tree for the selection of the experimental apparatus is shown in figure 2. Budget also plays a significant role in the selection of the IR camera since cooled systems are typically in the price range above a couple of ten thousand Euro, while noncooled systems, depending on the other specifications such as sensor resolution and lenses, are available at a fraction of the price (still typically around several thousand Euro). The cooled systems, operating at low IR spectra ranges, are more flexible, since they are suitable for operation at low and high temperature. The expected reaction conditions will strongly influence the choice of the materials and of the dimension of the reaction cell and the required type of IR camera to utilize. In figure 1, we underline the curve at 300 °C, which is the optimal temperature for our target application (Sabatier reaction). At this temperature, we can observe that the maximum of the spectral emittance curve lies around 6-7  $\mu m$ . The IR camera must be thus selected according to this fundamental parameter. In fact, the commercially available IR cameras are equipped with one among various possible detectors, which are sensitive to different IR wavelengths. Some of the available detectors are reported in table 1. The micro bolometer and the indium antimonite (InSb) detectors are suitable for applications up to 350 °C. Hence, their applicability in catalysis is limited to reactions occurring at low temperature, or with limited heat production. The QWIP (quantum-well infrared photodetector) is instead suitable for the investigation of emissions at lower wavelength, which means higher temperature. Therefore, this detector is the best choice for the investigation of reactions requiring high temperature. Furthermore, the QWIP detector allows recording a larger number of frames per second compared to the other systems presented. For catalysis applications, a stationary IR camera is generally sufficient to provide satisfactory results. Furthermore, in most of the practical situations investigated the use of cooled cameras is not required. However, these latter cameras undoubtedly show a better performance, simplifying the data collection.

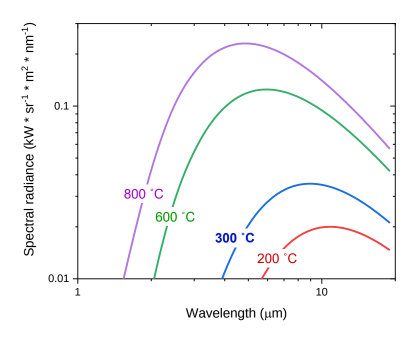


Figure 1 Determination of the wavelength of highest IR emission at different temperatures according to the black body emission spectrum

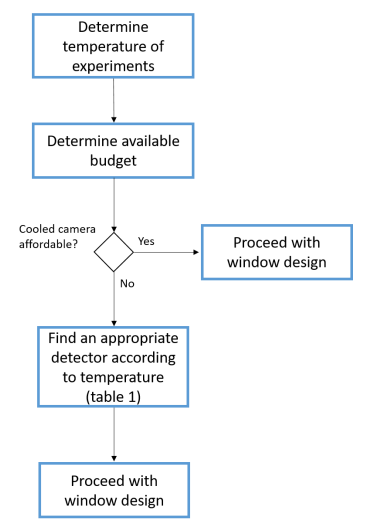


Figure 2 Decision tree for the selection of and IR camera for catalytic reaction investigation

Detector type	Temperature resolution (°C)	Temperature range (°C)	Exposure time (ms)	Frame rate (fps)	Spectral detection range (µm)
Microbolometer	0.075	-20 to +120 0 to +350	12	60	Roughly 7-17
Indium antimonide (InSb)	0.020	-10 to 55 50 to 150 150 to 350	1	170	Roughly 3-5
QWIP	0.025	-10 to +80 150 °C to 600 350 to 1500	16	380	Roughly 6-8

Table 1 characteristics of the main detector types employed in commercial IR thermography cameras [28–30].

# 2.2. Design of the reaction cell

The core of a system dedicated to the *operando* investigation of chemical reactions is the reaction cell. Once the IR camera has been selected, an appropriate system for the performance of the target reaction must be designed. Main elements to be considered are the window, the heating/cooling system and the shape of the reaction cell. These elements will be analyzed separately in the following subsections.

#### 2.2.1. Selection of the cell window

As the target of thermography experiments is the investigation of the IR emission of a catalyst, the reaction cell must be equipped with an IR at least partially transparent window. The window is selected mainly based on the spectral detection range of the camera and the reaction conditions — or vice versa. An overwiew on IR transmitting materials is listed in Table 2.

Every material listed is almost completely transparent for radiation with low wavelength (i.e. below 5  $\mu m$ ). At higher wavelength, the transmittance of the material can decrease rapidly with the wavelength, until becoming absorbant for lower frequency IR wavelenths and therefore no longer suitable for IR investigations. The transmission plot for BaF<sub>2</sub> (material selected for our case study) is reported in figure 3. We can observe that the positioning of the decreasing curve is also influenced by the thickness of the glass. Ticker windows become intransparent at lower wavelengths, as the transmission path is longer. However, the window must withstand the process pressure; hence, the glass must have a minimum thickness. The window thickness required to withstand a certain pressure can be calculated from the properties of the material and from the reaction pressure as follows:

$$PF = R^2 \times SF \times \Delta P \left[ Pa \cdot m^2 \right] \tag{2}$$

$$T = \sqrt{\frac{PF}{RM}} \quad [m] \tag{3}$$

Where PF is the pressure factor, R is the glass radius, SF is the safety factor (in our case 4),  $\Delta P$  is the pressure difference between the two sides of the glass and RM is the module of rupture of the material. The properties required for the calculation for various window materials are reported in

satisfactory solution.

table 2. The knowledge of the required material thickness, together with the information on the reaction temperature, allows for the final selection of the window material.

According to the principles described above, we derived a systematic decision tree for the determination of the ideal window material for an IR thermography reaction cell, which we report in figure 4. Firstly, temperature and pressure of the reactive phenomenon to investigate are determined. These are the key parameters for any further decision. Depending on the reaction temperature, the wavelength of the emitted IR radiation can be determined and therefore a first candidate material can be selected, among the possible IR transmitting materials of table 3. The information on material properties, together with the target process pressure, allow the calculation of the necessary window thickness to safely operate the IR thermography experiments. The combination of window thickness and IR wavelength are then utilized to determine the expected transmittance of the reaction cell at the target temperature. If a sufficient fraction of IR radiation is transmitted outside the reaction cell, the material is suitable for the thermography experiments and the system can be designed following the guidelines of the next section. If the transmission rate is not sufficient, the calculations must be repeated with another candidate material, until reaching a

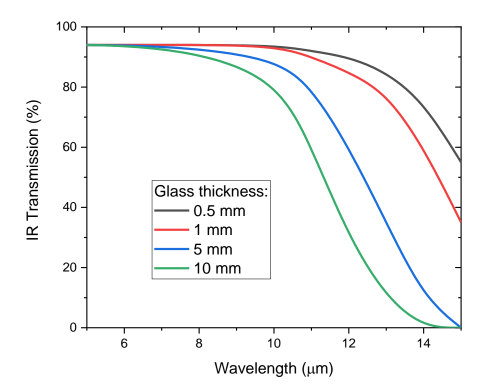


Figure 3 the IR transmittance of barium fluoride as a function of wavelength and thickness of the window (elaborated from [31]).

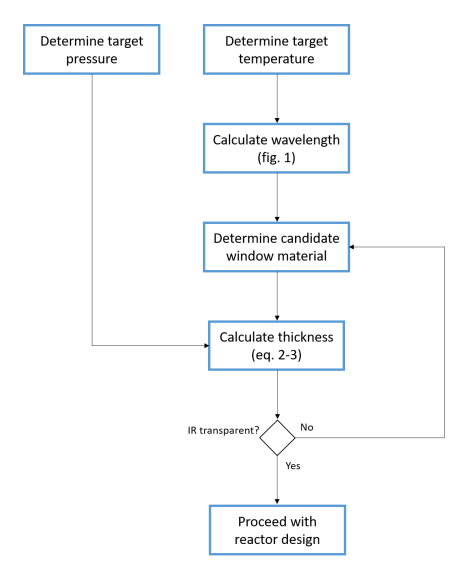


Figure 4 Decision tree for the determination of the appropriate material for the IR cell

Table 2 Properties of various IR transmitting materials (transmitting wavelength elaborated from [32], transmission at 8  $\mu$ m from [33])

Material	Transmitting wavelength range ( $\mu m$ )	Modulus of rupture (MPa)	Calculated required thickness for $\Delta P = 1$ bar (mm)	Transmission at 8 µm
BaF <sub>2</sub>	0.2-16	26.9	4.0	90%
CsI	0.6-30	5.6	7.1	45 %
CaF <sub>2</sub>	0.2-8	36.5	2.8	95 %
Fused silica	0.3-2	50	2.4	1 %
Ge	3-14	72.4	2.0	45 %
KBr	0.3-26	1.1	16.1	90 %
KCI	0.5-15	2.3	11.1	95 %
LiF	0.2-6	11.2	5.1	45 %
MgF <sub>2</sub>	0.15-8	49.6	2.4	75 %
NaCl	0.3-18	2.4	10.9	90 %
Sapphire	0.3-6	448	0.8	1 %

Si	2-10	125	1.5	1 %
ZnSe (used in	0.5-16	55	2.3	60%
this work)				
GaAS	2-18	71.9	2.9	55%
(material of				
the lens used)				

# 2.2.2. Selection of the heating system

In most of the practical uses of IR thermography for catalytic investigations, the reaction is conducted in a laboratory environment, using diluted gas. These conditions are necessary to avoid the influence of mass and heat transfer phenomena on the experimental results. For this reason, rarely the reaction operates in a self-sustaining way and an appropriate heating for the reaction cell is necessary. Furthermore, the external heating allows testing the catalyst at various temperature, to recover kinetic information. However, the installation of an appropriate heating system is challenging in the case of IR thermography experiments since the system must allow the clear distinction between the temperature changes due to heating and the effects of the chemical reaction. We designed a reaction cell in which heat is provided by a heating cartridge placed immediately below the reaction plane. In this way, the entire reaction cell is heated at the same temperature thanks to the rapid heat transfer through the metal walls. Furthermore, the heat is directly transferred to the catalyst, which quickly reaches the target temperature upon activation of the heating system. According to the dimension of the reaction cell, it may be convenient to install more than one heating cartridge, to ensure a uniform temperature (circular heating plates or multiple heating cartridges). During the experiments, we observed that special attention had to be posed to the control system of the heating system. In fact, the standard temperature control of the heating cartridge (feedback loop controller) tends to compensate for the temperature increase due to the chemical reaction, thus strongly interfering with the experimental observations. To avoid this feedback loop, the heat controller / heating system has to be set to a constant power output rather than a temperature set point. In this case, the aimed reaction base temperature has to be found manually with power set point iterations. In Figure 5, we show the result of a test run of the Sabatier reaction with a fixed power output set point for the heating cartridge. Some background noise in sinusoidal waveform is visible due to the external heating. This is likely due to the regulation of the power which is not constant but probably set via on/off in certain time steps which results in the reported waveform of the baseline. However, the wave amplitude is less than 1°C and therefore the baseline is suitable for experiments where the heat increase/decrease is clearly above 1°C from the baseline. In the specific case of figure 5, the interference of the background noise does not influence both the observation of a steady state temperature gradient (2.7 °C difference between reactor inlet and outlet) and a transient hotspot (ca. 1.8 °C for the point at the reactor outlet, P13 in figure 5). These results validate the appropriateness of the power set point method for the IR thermography experiments, showing that it is possible to achieve a limited interference between the reaction effects and the external heating. However, this creates a larger effort in the experimental design phase, as the system has to be calibrated in all the condition tested, to ensure that the right power is fed to the heating system and the target temperature is reached and maintained. Additional attention should be posed towards the formation of temperature gradients between the catalyst surface and the reactor window. In fact, the temperature outside of the window may be significantly

lower than the temperature of the catalytic bed, resulting in a temperature gradient within the height of the channel, which makes challenging the modeling of the heat transfer phenomena. A proposed solution to this issue involves the use of a heated blower to supply hot air on the external face of the glass, thus reducing the temperature gradients [24]. Otherwise, it is possible avoid this problem by installing the entire setup in a temperature-controlled environment (e.g. oven with an IR transmitting window).

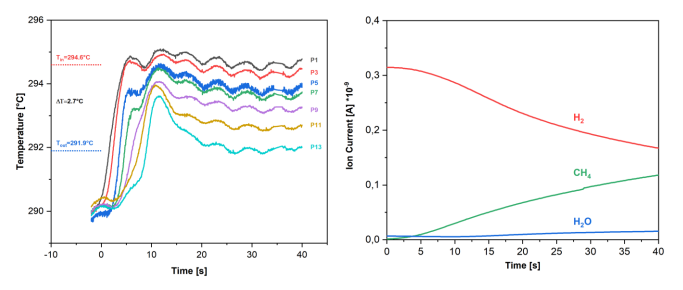


Figure 5 Temperature profiles measured with an IR camera during the Sabatier reaction over a 2% Ru/Al $_2$ O $_3$  catalyst.

#### 2.2.3. Coupled concentration measurements

In order to obtain a full picture of the chemical phenomena occurring during IR thermography experiments, it is necessary to couple the detailed temperature analysis with the determination of timely resolved concentration curves. These must be exactly timed with the IR temperature profiles to enable a correlation between temperature and concentration profiles. Considering the example of figure 5, we can observe that the transient hotspot recorded by the IR camera takes place in the first 20 seconds of the experiment. Therefore, the concentration profiles at the reactor outlet (and ideally also along the axial coordinate of the reactor) must be recovered with a similar frequency (several points during these 20 seconds). Due to this strict requirement, only few concentration analysis tools are suitable. Such a rapid sampling rate can be achieved only with spectroscopic methods, such as mass spectroscopy (MS) or Fourier transform infrared spectroscopy (FTIR). A standard method like gas chromatography (GC) offers instead a too low sampling rate to be coupled with IR thermography. In order to achieve a faster sampling rate, the measurement method can be optimized to include only information about the species of interest. This is the case of figure 5b, where the mass spectrometer has been employed only in the determination of the chemical compound that are consumed or formed in the Sabatier reaction (i.e. CO2, H2, H2O and CH4). In this way, the sampling rate can be significantly increased. For the specific case of figure 5b, 50 spectra per second were recorded. This frequency integrates well the IR thermography results, indicating that the initial hotspot is generated by water adsorption on the catalyst surface, as pointed out by the delay in the observation of water at the reactor outlet compared to the first appearance of methane. Additionally, the use of a spatially resolved technique, such as the analysis in multiple positions of the catalyst bed by use of a movable capillary, can integrate a large amount of

temperature data points with a large amount of concentration data points, creating an ideal dataset for kinetic modelling and model-based reactor optimization.

### 3. Design of the IR thermography experiments

IR camera, reactor window and heating system are the most critical aspects to be addressed when designing an experimental system for IR thermography measurements. In the previous sections, we defined a systematic procedure for the design of these elements and determined the characteristics that a system must contain to provide significant experimental results. However, a crucial point for the recovery of experimental evidences about transient and steady state chemical phenomena is the appropriate design of the thermography experiments, in order to avoid the influence of undesired side effects, caused by heat and mass transfer limitations or by the non-ideality of the system. In this section, we define some general guidelines to correctly design IR thermography experiments. These involve the determination of the optimal procedures to perform the selected chemical reaction, in terms of adaptation of the reaction kinetics and feed sequence of the reactants.

# 3.1. Formulation of heat balances

As IR thermography is aimed at the analysis of the temperature of solid and/or of the gas phase in a chemical reaction, the reaction conditions must be selected with care. In this sense, the most common reaction engineering tools are greatly helpful for the experimental design of IR thermography experiments. Primarily, a fundamental parameter to be determined for designing the experimental procedure is the catalyst emissivity. This parameter can be evaluated by performing calibration experiments with the IR camera, exposing the catalyst to an inert gas such as He. The most common calibration method involve the comparison of IR thermographs with the temperature measured by a thermocouple [23]. Emissivity is dependent on the temperature; therefore, a calibration curve must be determined by evaluating the results collected over the entire temperature range of interest. Detailed calibration curves may consider also the influence of focal distance and humidity [34].

Once the emissivity of the catalyst in relevant conditions is determined, the heat losses by convection and irradiation can be calculated via equation 3 & 4. The radiative heat flow is calculated as:

$$Q_r = \varepsilon \sigma A (T^4 - T_e^4) \tag{3}$$

Where  $\varepsilon$  is the emissivity,  $\sigma$  the Stefan-Boltzmann constant, A is the exchange surface, T the process temperature and  $T_e$  the external temperature.

The convective heat flow is calculated with formula 4:

$$Q_c = UA(T - T_e) \tag{4}$$

Where U is the heat transfer coefficient, obtained from appropriate correlations for the Nusselt number. In most of the micro reactors used for laboratory applications, the flow regime is laminar; therefore, the contribution of convective heat transfer is limited [35].

# 3.2. Reaction engineering considerations

The heat balance considers the heat losses by convection and irradiation (conduction can be neglected in almost all the practical cases) and the heat provided by reaction and external heating. These elements can be implemented in an appropriate reactor model to estimate the temperature profile on the reactor and verify whether IR thermography is a suitable method for the analysis of the reaction. For this preliminary assessment, a simple 1D pseudo-homogeneous model can be sufficient, providing some general information on the expected extent of the reaction hotspot.

#### Estimation of temperature profiles in the Sabatier reaction

In our case study, we used a 1D pseudo-homogeneous reactor model, simulating the Sabatier reaction over a 0.5% Ru/Al<sub>2</sub>O<sub>3</sub> catalyst (modelling details are reported in the supplementary information). The results of the simulations, which show the predicted steady state profiles, are reported in figure 6. Surprisingly, we observed that the reaction occurring on this catalyst would generate a limited reaction hotspot (ca. 1.5 °C) and that the CO₂ conversion would be distributed over the entire reaction length. Hence, in this reaction system, it would be difficult to distinguish the reaction-generated hotspot from the natural background noise originated from the heating system (see Figure 5a). Thermography experiments in this specific system would not allow a reliable determination of dynamic effects in the reaction, but the practically isothermal reactor would be an ideal solution for the kinetic analysis of the system since this would minimize the influence of nonideal conditions on the experimental results. In this latter case, the system should be complemented with spatially resolved concentration profiles, to obtain a large and consistent dataset of temperature and partial pressure measurements for the kinetic modelling phase. In order to design a system suitable for the experimental analysis of transient phenomena, the heat production of the system should be increased or the heat losses should be decreased. In our case, we decided to increase the heat production of the system by using a catalyst with higher Ru load. The red and dashed curves of figure 6 show the calculated temperature and CO<sub>2</sub> conversion profiles for the 2% Ru/Al<sub>2</sub>O<sub>3</sub> catalyst. We can observe that the Sabatier reaction is faster on this catalyst, resulting in a different distribution of the heat production. Most of the CO<sub>2</sub> conversion occur in the first third of the reaction, producing a remarkable hotspot at the reactor inlet (ca. 5.5 °C). In these conditions, it is possible to distinguish between the heat originated from the reaction and the contribution of the heating system. We expose the experimental validation of these calculations in section 4. According to these simulations, this reactive system is ideal for the determination of the dynamic effects of the Sabatier reaction. On the contrary, the presence of a remarkable hotspot in this configuration makes the system unsuitable for the kinetic analysis of the system. If the target of the study is the kinetic modelling of the reaction over the 2% Ru/Al<sub>2</sub>O<sub>3</sub> catalyst, this hotspot should be removed, for example by diluting the catalyst with inert material. Another possibility would involve the dilution of reactant with inert gas, but the influence of this addition on IR transmittance must be verified to avoid undesired effects.

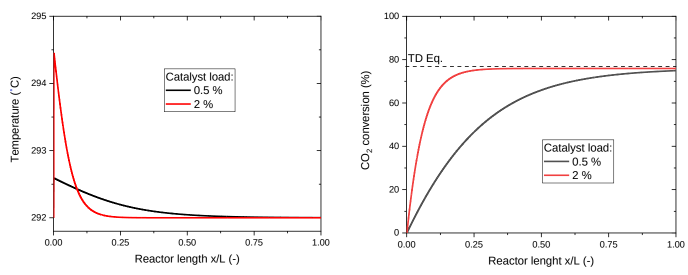


Figure 6 The predicted a) temperature profile and b)  $CO_2$  conversion for the Sabatier reaction over a 0.5% (black line) and 2% (red line)  $Ru/Al_2O_3$  catalyst

# 3.3. Experimental design routine for IR thermography tests

This initial screening is fundamental to determine the suitability of an experimental setup and of a catalytic system for the target application. Even though the preliminary results are not sufficient for a detailed formal analysis of the system, they can significantly increase the probability of success in the IR thermography experiments. In particular, the determination of an appropriate experimental design can decrease the number of experiments required for the target application, allowing for significant cost savings. On the base of the considerations exposed in this section, we developed a decision three for the experimental design of IR thermography tests, which is reported in figure 7. The initial point of the experimental design follows the determination of the characteristics of the IR thermography hardware, as described in section 2 (IR camera and IR cell). These tools fix the most important properties for the determination of the heat transfer properties, such as transmissivity and gas flow ranges. On the base of the hardware available, it is possible to select a first candidate catalytic material for the experiments. With the catalyst and reactor properties, it is possible to operate a first approximate simulation of the reaction in the IR cell. This allows the determination of the extension of the reaction hotspot, which is the main parameter to consider for further experimental design. According to the target of the experiments, an isothermal or a non-isothermal reactor may be needed.

# Determination of reactor requirements

In the former case, which is typical of kinetic modelling experiments, the hotspot should be limited, not to damage the experimental results with parasitic effects present in the hotspot. The critical threshold for the system can be set to a temperature variation in the hotspot equal to 1 K. This temperature is chosen as it is a typical oscillation of a common heating system (cfr. Section 2.2.2). If the calculated extension of the hotspot is larger than 1 K, it is appropriate to take measures to decrease this phenomenon, for example by diluting the catalyst or the reactive gas flow. In the case of non-isothermal experiments, the situation is reversed. If the calculated hotspot temperature is below 1 K, it will be necessary to increase the heat production of the system, to ensure that the phenomena to investigate are well visible and distinguished. In this case, this can be achieved by increasing the reaction rate (e.g. by using a catalyst with higher load of active phase) or by

decreasing the heat losses (e.g. by decreasing the temperature difference between the two sides of the glass window). Upon verification of all the conditions in the decision tree, the assumptions can be validated by appropriate IR thermography experiments. In figure 8, a procedure for the realization of these tests is elucidated. IR cameras can determine the temperature of every single pixel in its focal view, but not all these points have a relevance for the analysis. Hence, only the necessary data points should be collected through the IR camera acquisition software. In the case of panel 1 of figure 8, we selected 13 equally spaced points on the middle axis of the reactor. This is a reasonable in most of the laboratory experiments, as the axis of the reactor is the stationary point of the Poiseuille curve of laminar flow in the reactor and therefore the location with lowest influence of the borders of the reactors. The selection of these limited amount of data points facilitates the analysis and diminish the need for storage, without decreasing the significance of the experimental dataset. Only in specific applications, such as the validation of CFD calculations, the entire focal view can be of interest. Once the measurement points are selected, the experiment can be performed and the temperature field is recorded (panel 2 of figure 8). In the figure, the presence of a Poiseuille curve for laminar flow during the experiment is evident, confirming the need for recording the data points on the middle axis of the reactor (see Section 4.2). An appropriate time spacing of the recordings should be defined, in order to compromise between the significance of the data and the amount of data recorded. For fast dynamic experiments, an interval of 1 second or lower is suitable. However, for phenomena occurring over large time scales, recording times up to minutes can be appropriate. This is the case, for example, of the analysis of catalyst deactivation [9]. As a last step of the analysis, the temperature data are integrated with appropriate concentration measurements (figure 8, panel 3). As mentioned in section 2.2.3, these concentration profiles must be consistent with the temperature measurements and may determine the time scale of the analysis. The coupling of these two analysis opens the way to the detailed investigation of chemical phenomena, with large resolution in time a space. The power of this experimental tool is elucidated in the following section with some specific case studies, concerning both dynamic and steady state catalytic processes.

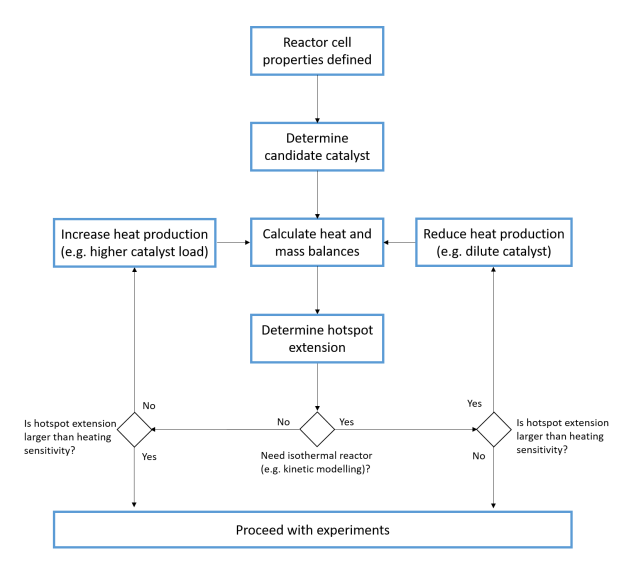


Figure 7 Decision tree for the determination of an appropriate experimental design methodology for IR thermography experiments

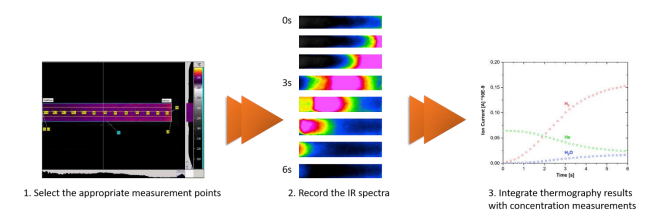


Figure 8 Routine for the successful realization of IR thermography experiments: the routine starts with (1) the selection of appropriate measurement points, continues (2) with the recording of the IR spectra, which have to be integrated with (3) sufficient concentration measurements.

# 4. Examples of IR thermography experiments

This section is dedicated to the elucidation of selected experimental results obtained with the methodology presented in the previous sections. The complete dataset is accessible in our previous work [24]. The goal of this section is the validation of the methodology with relevant experimental results. Therefore, we focus on specific aspects of the experimental results that elucidate the importance of an appropriate design of the experimental setup in the achievement of significant results.

## 4.1. Experimental setup

Figure 9 shows the flow diagram of the experimental setup and a picture of the installed reaction cell together with the IR camera. The reaction cell is manufactured from aluminum and is equipped with a zinc selenide window. The catalyst bed is placed below this window. A round heating cartridge (Maxiwatt, Spain) with a maximal power of 250 W is placed below the catalyst bed. A Jumo Imago 500 controller adapts the power of the heating cartridge to ensure a stable and constant power input. The IR camera used in the experiments is a VarioCAM hr from InfraTec (Germany). The camera is operated through the software IRBIS 3.1 PRO, directly provided by the camera supplier. Calibration of the camera was performed in He flow, by adapting the reaction cell temperature between 25 and 280 °C and comparing the measurements with the value recorded by a K-type thermocouple placed below the catalyst bed. please refer our previous publication [24]. The catalyst employed in the measurements is a 2% Ru/Al<sub>2</sub>O<sub>3</sub>, whose measured emissivity is 0.85. The reaction cell is loaded with 172 mg of the catalyst. The catalyst is prepared by incipient wetness impregnation following the recipe elucidated in [36]. For a complete overview of the experimental setup, and for further details on the catalyst characterization, please refer to our previous study [24]. The reaction products are analyzed by means of mass spectroscopy (MS). The MS instrument is a Pfeiffer Omnistar GSD320. In order to achieve a fast sampling speed (20 ms per spectrum), the MS analysis is limited to the relevant mass for the Sabatier reaction experiments, referring to CH<sub>4</sub>, CO<sub>2</sub>, CO and H<sub>2</sub> (i.e. m/z 2, 4, 15, 18, 28, 44). The absence of other species in the product stream was verified in preliminary experiments. The procedure for quantitative analysis of the MS spectra through the Faraday detector follows our previously developed methodology [37].

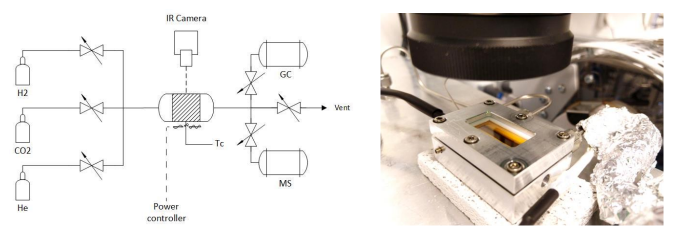


Figure 9 a) schematic representation of the experimental setup; b) a picture of the reaction cell in the focal view of the IR camera

# 4.2. Surface reduction experiments

When exposed to an oxidizing atmosphere (e.g. during storage), the  $Ru/Al_2O_3$  catalyst tends to be superficially oxidized. Therefore, prior to any further experiment, the metallic composition of the catalyst must be recovered by reacting the surface oxygen with  $H_2$ , following the reaction:

$$RuO^* + H_2 \leftrightarrow Ru + H_2O \qquad \Delta H_0^R \approx 482 \, kJ/mol$$
 (5)

This reaction is highly exothermal and takes place until full reduction of the catalytic surface. A specific procedure for the surface reduction of the catalyst was developed in our previous studies [18,19]. In general, a temperature programmed surface reaction (TPR) is operated to completely remove the oxygen from the catalytic surface, heating the sample from room temperature to 700 °C. However, the reaction takes mainly place in a limited temperature interval, which can be accurately determined thanks to IR thermography. In fact, reaction (5) is fast and exothermal, so that its occurrence is easy to detect by recording the temperature change on the catalyst surface, but the extension of the reaction can be observed only with an instrument with sufficiently high sampling rate. Thus, the temperature of activation of the surface reduction can be determined by recording with the IR camera the TPR experiment. Preliminary experiments showed that, for the catalyst of this study exposed to the atmosphere, the H₂ reduction activation temperature is ca. 290 °C. Hence, we set the baseline temperature of the reaction cell loaded with the oxidized catalyst at this temperature and we fed 10 mL/min of H<sub>2</sub>. The raw IR thermography data recorded during the surface reduction are shown in the second panel of figure 8. The recorded temperature at the center of the reactor on the middle axis are displayed in figure 10a. The temperature used for further elaboration is recorded on the axis of the reaction in order to minimize the influence of wall effects from the laminar flow, which are evident in the full raw IR data. The corresponding MS spectra are shown in figure 10b. The complete phenomenon takes place in ca. 7.5 s. During this time, the temperature measured is higher than the baseline temperature. Initially, the temperature rises from the baseline due to the activation of the reaction, showing an initial thermal runaway. When the oxygen on the surface starts depleting, the reaction rate decreases, forming a maximum in the recorded temperature. The extent of the temperature hotspot is ca. 14.6 °C. The temperature decreases then quickly, for the rapid deactivation of the reaction due to full consumption of oxygen. The comparison with the MS spectra recorded at the reactor outlet shows a good agreement with the thermography measurements. In fact, the MS spectra show a progressive increase in the production of H<sub>2</sub>O, with a clear flex after ca. 7 seconds. The IR spectra reach steady state conditions after 20 seconds. This longer time to steady state is due to the presence of dead volumes between the reactor and the spectrometer and reflects the intrinsic averaging occurring in the MS. In fact, while the temperature is recorded exclusively on the reactor axis, the gas concentration is averaged on the entire cross section of the reactor. Therefore, the resulting concentration profile contains particles that underwent different residence times in the reactor and should be considered with care to avoid misinterpretation of the results. This example shows how the different resolution between IR thermography and other analytical techniques challenges the interpretation of results. For example, the use temperature data taken from a punctual location in the reaction may not agree with integral concentration data due to the non-ideality of the system. However, the high resolution of IR thermography is a fundamental help in the characterization of the transient phenomena. Further elaboration of the combined temperature and concentration profile can open the way to the determination of further properties of the system. As an example, by combining the information on

the duration of the reaction and on H<sub>2</sub> conversion, it was possible to calculate the surface coverage of the catalyst with oxygen, which was quantified as a single layer [24].

The use of IR thermography in the study of dynamic processes evolving towards a steady state can reveal more insights both in the way the system approaches the steady state and in the properties of the steady state operation. In order to elucidate the challenges and opportunities in the study of these kind of processes, we report in the next section some exemplary results from the study of the Sabatier reaction.

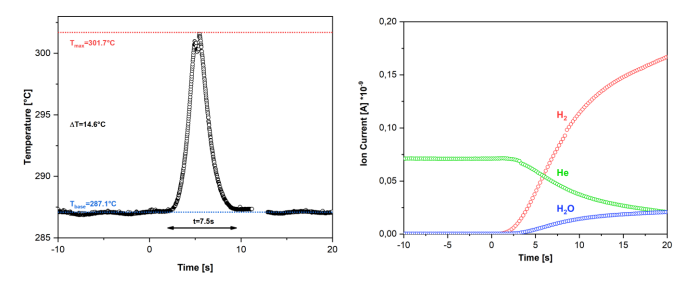


Figure 10 a) the temperature profile recorded in the center of the reaction cell during the reduction of the oxidized 2%  $Ru/Al_2O_3$  in  $H_2$  flow; b) evolution of the MS spectra during the experiment. Experimental conditions:  $H_2$  flow: 10 mL/min, P=1 bar, baseline temperature = 287 °C.

# 4.3. Sabatier reaction experiments

The analysis of the Sabatier reaction shows a double level of challenge, as the reactive phenomena involve both transient and steady state temperature variations. Figure 11 shows the steady state temperature profile in the reactor, as well as the Ion currents recorded by the MS instrument over time. We can observe that, in the MS spectra, steady state conditions are reached after ca. 150 s. After this time, the reaction occurs in the reactor showing the characteristic temperature hotspot at the reactor inlet. The steady state concentration and temperature profiles are determined by the interplay of kinetics and thermodynamics [21]. In the lab-scale reaction cell, the extension of this hotspot is limited (ca. 2.5 °C), but it is revealed well by the IR camera. As explained in section 3.2, the reaction cell was designed with the purpose of making this phenomenon visible. The experimental results are in good agreement with the preliminary modelling results and show the importance of an appropriate model based experimental design to perform the experiments in a proper way and with the appropriate testing equipment. The quick change in the steady state temperature over the axial coordinate, in particular the closeness of the hotspot to the reactor inlet, makes the determination of the temperature profile by means of thermocouples quite challenging. In fact, even though a moving thermocouple is placed inside the reactor, the position of the hotspot remains difficult to locate and the presence of the thermocouple can influence the reaction. Furthermore, it may be difficult to distinguish the gas and the surface temperature [38]. Therefore, we demonstrate here that the use of IR thermography is an optimal solution for the study of highly exothermal reactions and for the recovery of useful kinetic information to be used in model based reactor design and

optimization routines [9,39]. The required effort for the design of an appropriate reaction cell is compensated by the high quality of the obtained experimental data and can be minimized by the routines presented in this study.

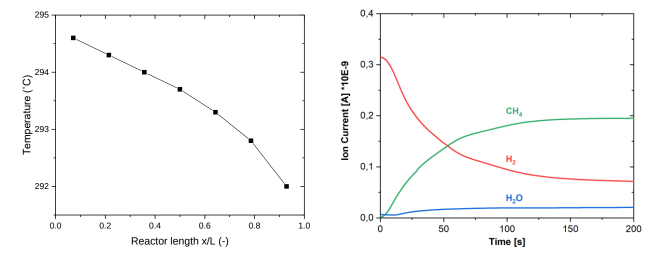


Figure 11 a) axial temperature profile at steady state and b) concentration profile over time in the Sabatier reaction. Experimental conditions: total gas flow: 20 mL/min,  $H_2:CO_2 = 4:1$ , P=1 bar, baseline temperature =  $290 \, ^{\circ}\text{C}$ .

In addition to the steady state results, the designed experimental setup allows the observation of the Sabatier reaction during the initial transient. The evolution of the temperature profile upon feed of a stoichiometric H<sub>2</sub>/CO<sub>2</sub> mixture over the reduced catalyst is shown in figure 12. The reaction starts immediately upon injection, as stated by the temperature profile after 2.5 seconds. Initially, only a small fraction of the reactor is involved in the reaction, because the remaining part is still filled with inert gas. When the inert gas is replaced on the entire reactor length, we observe that the temperature increases progressively, due to the course of the reaction heating up the catalyst. The first three measurement points considered reach quickly the steady state conditions, while the rest of the reactor undergoes the formation of a transient hotspot. We observe that the entire catalyst bed reaches a temperature above 294 °C after 10 seconds and that the temperature at the end of the reactor progressively decreases afterwards. This progressive cooling of the reactor evolves towards the steady state temperature profile afterwards until reaching the steady state conditions, as described above and in figure 11a. An explanation of this phenomenon can be found by integrating the temperature profiles with the information on product concentration from figure 11b. The catalyst is initially fully reduced, therefore no oxygen containing species are present on the surface. Hence, upon feed of the stoichiometric H<sub>2</sub>/CO<sub>2</sub> mixture, the local H<sub>2</sub> concentration results over-stoichiometric. This shifts the thermodynamic equilibrium, allowing for a higher conversion than in steady state conditions. In this phase, all the O<sub>2</sub> fed in the form of CO<sub>2</sub> is converted into oxygenated surface species and no H<sub>2</sub>O product is formed, as visible in figure 11b (first 10-20 seconds of the reaction). When the local equilibrium between reduced and oxidized surface species is reached (details about the nature of these species can be found in our previous studies [40,41]), the reaction reaches the steady state conditions, with the characteristics described above. These examples shows how the effective coupling of IR thermography and fast concentration measurements (e.g. by means of MS) allows the efficient performance of transient and steady state experiments, recovering a large amount of high-quality experimental data with low experimental effort (i.e. minimizing the need for manual interaction with the system). Furthermore, the raw data contain a much larger amount of points in the time and space scale, which can be further elaborated later, if required. Additionally, as important advantage compared to standard temperature measurement techniques, the position of the analyzed measurement point can be changed anytime, increasing the robustness of the experimental procedure and reducing the requirement of repeated measurements, thus minimizing the effort for the experimental campaign.

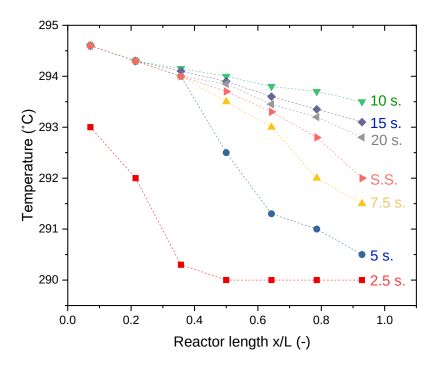


Figure 12 Evolution of the surface temperature over time upon feed of  $H_2$  and  $CO_2$ . Experimental conditions: total gas flow: 20 mL/min, H2:CO2 = 4:1, P=1 bar, baseline temperature = 290 °C. (S.S. = Steady State)

#### 5. Conclusions

In this paper, we derived a systematic methodology for the use of IR thermography in chemical reaction engineering. We defined the requirements of an experimental system to appropriately record the chemical phenomena and to avoid the influence of external occurrences on the measurements. To this scope, we elucidated the characteristics necessary for the reaction cell used in the experiments and the ideal experimental conditions to achieve during the measurements. Furthermore, we provided selected examples of IR thermography experiments to investigate the Sabatier reaction and the lessons learned from these experiments.

IR thermography appears as an ideal candidate for the advancement of catalysis and chemical reaction engineering, thanks to the intrinsic capability of providing new insights thanks to the high resolution both in space and in time. In this sense, it can give an important contribution towards development of high-throughput experimental techniques. However, in this paper we observed that the implementation of the technique is challenging and some aspects in the design of the experimental setup must be correctly addressed:

 The IR camera must be chosen in agreement with the temperature of the application and appropriately calibrated. Most of the IR thermography applications in catalysis require that the camera can follow a large temperature window without incurring in signal saturation problems.

- The experimental setup must fulfill both requirements of IR transparency and withstand the process pressure. As a tradeoff exists between IR transmittance and thickness of the glass, the appropriate material must be selected. Several materials are appropriate to this scope and should be screened with the routines derived in section 2 to find the most suitable.
- The reaction cell must be heated with the appropriate system (i.e. in power control) to
  avoid contamination of the experimental results by external factors. Additionally, particular
  attention should be posed to the insulation of the system to avoid the formation of
  excessive temperature gradients between the two sides of the window. Several heating
  methods such as cartridge heaters, heating plates and IR transmitting ovens were discussed.

Furthermore, the characteristics of the reaction cell must match the requirements of the reactive system investigated, that is:

- The system must be able to record the effects of the chemical reaction on the catalyst temperature. This means that the oscillation of the temperature baseline must not cover the hotspot of the chemical reaction and the latter must be sufficient to compensate for the heat loss by convection and radiation.
- The system can be optimized prior to the investigation with a model based experimental design routine, so that the experimental effort is minimized by avoiding the trial and error approach. In this sense, we proposed a model-based experimental design routine in section 3, validated with the case studies presented in this paper.

We observed that, with the help of IR thermography, the investigation of exothermic reactions, such as the Sabatier reaction, could be performed with an important reduction of the experimental effort. This is possible because large datasets of transient and steady state temperature measurements can be obtained from a single experiment, if the system and the experiments are designed appropriately. Hence, the amount of resources required for the preparation of an experimental dataset at the base of a kinetic modelling routine can be substantially reduced and the quality of the data used for model validation and optimization purposes can be greatly improved thanks to the high time and space resolution. We thus expect that IR thermography will play an important role in the advancement of catalysis and chemical reaction engineering in the future, guiding the development of new experimental and numerical methodologies for laboratory and industrial practice.

# Acknowledgments

This work did not receive any specific funding. We thank the organizers of the EFCATS Summer School 2020 for the invitation to contribute to this special issue and the Lib4Ri library of the research institutes of the ETH Domain (CH) for the support to publish open access.

# **CRediT authorship contribution statement**

**Robin Mutschler:** Conceptualization, Methodology, Investigation, Formal analysis, Writing - Review & Editing. **Emanuele Moioli:** Conceptualization, Investigation, Formal analysis, Data curation, Writing - original draft, Resources, Funding acquisition.

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