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An overview of differential scanning calorimetry

$I = I - I$

In differential scanning calorimetry (DSC) the difference in power $(\Delta P, \text{ in } mW)$ required to heat a sample (S) and an inert reference (R) is determined as a function of temperature (T). There are many commercially available DSC instruments, the designs of which usually vary only in terms of furnace arrangement and/or the number and positioning of thermocouples. Early instrument designs plotted the difference in temperature between S and R (differential thermal analysis, DTA) but this term is no longer in common use. DSC instruments are compact, with a footprint typically no larger than 0.5 \times 0.5 m².

DSC is one of a group of techniques that make measurements at controlled temperature. Collectively these are known by the term thermoanalytical and the field is termed thermal analysis (TA).¹ The International Confederation of Thermal Analysis and Calorimetry (ICTAC) degnes the nomenclature and calibration methods used throughout the TA f eld.

$\mathbf{E}(\mathbf{r})$ and a point application

DSC has widespread application in many diverse \hat{p} elds because it does not require the sample material to possess any specific functional or chemical property, save that it changes its heat content when undergoing a phase change. The small size of the sample pans limits the sample mass to ca. 5–10 mg and so solids are usually studied. Large volume instruments are available for solutions. The instrument can detect thermally-driven phase transitions (such as melting, crystallisation and glass transitions) as well as the loss of volatile components. It is possible to purchase DSC instruments capable of studying solutions, but these have larger cells and operate at slower heating rates, and so are not considered in this report.

Key features All DSC instruments control temperature (T) with respect to time (t). The heating rate (b) can be:

– Linear $(dT/dt = b)$

– Modulated (e.g. $dT/dt = \sin b$)

– Stepped (e.g. a number of isothermal periods at discrete temperatures)

– Sample-controlled (e.g. the sample response governs b).

All DSC instruments use a heater (termed a furnace) to supply power to the sample and reference materials. Where a common (or single) furnace is used to heat S and R, the instrument is of a heat- $flux$ design and where separate furnaces are used to heat S and R, the instrument is of a power-compensation design. Knowledge of the arrangement is important. Heat- ux instruments tend to have a greater thermal mass and

so usually have lower maximum heating rates (typically up to ca. 200 °C min⁻¹). Power-compensation instruments, having smaller thermal mass furnaces, can achieve faster heating rates (typically up to ca. 750 °C min⁻¹). Solid-state (i.e. etched silicon chip) calorimeters, possessing very little thermal mass, can achieve heating rates of the order of 10^6 °C s⁻¹. Similarly, measurements may be performed at denned cooling rates. The cooling rates achievable will be dependent upon the cooling system (typically either a refrigerated cooling system (RCS) or a liquid-nitrogen circulator). Again, the maximum rates attainable will be fastest for solid-state systems and slowest for heatux instruments. A power-compensation instrument equipped

with an RCS would typically cool to -60 °C at -100 °C min⁻¹.

$B = \frac{1}{2}$ basic principles

Sample and reference materials are sealed in pans (or crucibles) and placed in the instrument before being heated (or cooled) in accordance with a user-defined programme. Pans are typically made of pressed aluminium (for experiments up to $600 °C$) or aluminium oxide or gold (if higher experimental temperatures are required). Stainless steel, gold-plated stainless steel highpressure crucibles and/or sealed glass crucibles are used for decomposition studies at high temperatures.

 \mathbf{H} - \mathbf{f} . In heat-ux DSC, a common furnace heats the sample and reference pans and the temperature difference (ΔT) between them is recorded, usually with thermocouples placed close to, but not in contact with, the pans. The number and positioning of the thermocouples will vary between instrument designs. The power change occurring in the sample is directly proportional to the temperature difference and is calculated by multiplying the data by a constant of proportionality, determined by calibration with a certiped reference material (CRM). Power compensation DSC. In power-compensation DSC, separate furnaces heat the sample and reference materials. The instrument varies the power supplied by the two furnaces to maintain the temperature difference between the sample and reference at a constant value (usually close, but not equal to, zero). The power difference (ΔP) between the sample and reference is thus measured directly.

$D = -1$ measurements of $T = 1$

DSC measurements comprise contributions from two sources; heat capacity (C_p) effects and any other processes (phase transformations or chemical reactions) that the sample might undergo (represented by the generic term f(T,t)).

$$
\frac{d\mathcal{V}}{d} = \rho \frac{d}{d} + \rho \left(\rho, \rho \right) \tag{1}
$$

If the sample undergoes a phase change or chemical process, or there is a change in heat capacity, there will be a concomitant event in the DSC data. Typical DSC data, in this case an endotherm resulting from melting of a pure compound, are shown in Fig. 1. It should be noted that endotherms may be plotted positive or negative depending upon whether the instrument calculates ΔP as S–R or R–S. The convention varies between

manufacturers and so the direction should be indicated on the y-axis. Several parameters may be determined, including the heat of fusion (AH_{fus})

proceed in the forward direction only and so are termed nonreversing. Since the kinetic response follows a sine function it should be 90° out of phase with the heat capacity term.

The utility of MTDSC comes in being able to deconvolute the overall heat- ow signal into these two components. Deconvolution requires calculation of the underlying heat ow signal (which is the average of the modulated response – equivalent to the heat ow signal that would be recorded for a normal DSC experiment performed at the same underlying linear heating rate). This can be considered equal to:

Underlying heat flow =
$$
_{p}b + \frac{1}{2}
$$
 (1) (4)

Heat capacity re ects the rise in temperature for a given input of heat and so can be determined by comparing the amplitude of the modulated heat- ow (A_{mhf}) with the amplitude of the modulated heating rate (A_{mhr}) :

$$
p = \frac{mhf}{mhr}
$$
 (5)

The amplitudes are determined using a Fourier Transform. From eqn (4) and (5) it can be seen that the heat capacity (reversing) component of the underlying heat ow is given by:

$$
Reversing heat flow = b \frac{mhf}{mhr}
$$
 (6)

Hence the kinetic response (non-reversing) component of the heat ow can be determined by difference:

Non-reversing heat-flow = Underlying heat flow – b
$$
\frac{\text{mhf}}{\text{mhr}}
$$
 (7)

Heat ow data recorded with MTDSC thus allows separation of processes into reversing or non-reversing events. This aids both identiacation and isolation (if multiple events occur at the same temperature) of processes, although being a mathematical routine it is possible to introduce artifacts into the reversing and non-reversing data if the modulation parameters are not selected carefully.

 $E = \frac{1}{2}$ and $E = \frac{1}{2}$ and $E = \frac{1}{2}$. Pans comprise a base and a lid. The seal between pan and lid can be hermetic (air-tight) or non-hermetic and correct selection is important. If the sample contains water that will evaporate into the headspace of the pan, different responses will be seen in hermetic and non-hermetic pans (the headspace of a hermetic pan will become saturated and evaporation will stop). Similarly, if there is a large expansion in volume of a sample (as it melts for instance) the increase in pressure inside a hermetic pan can cause the seal to fail (although hermetically sealed high-pressure pans can cope with up to 250 MPa pressure). For this reason, some pans have a pinhole in the lid to prevent a build up of pressure. In all cases, the sample and reference pans should be as closely matched in weight as possible.

 $\mathbb{E}_{\mathbb{E}_{\mathbb{E}_{\mathbb{E}_{\mathbb{E}_{\mathbb{E}}}}\left| \mathbb{E}_{\mathbb{E}_{\mathbb{E}}\left[\mathbb{E}_{\mathbb{E}_{\mathbb{E}}}\right] \right|}}$. The most important parameter is the heating rate. DSC heating rates typically range between 2– 200 °C min⁻¹. Varying heating rate can help identify whether thermal transitions are thermodynamic or kinetic in nature, because kinetic events show a dependence on heating rate. It is thus good practice to repeat measurements at two heating rates, an order of magnitude apart (2 and 20 or 20 and 200 \degree C min⁻¹ for instance). It is also advisable to cool the sample after the $\hat{\textbf{r}}$ arst heating run and then reheat it using the same parameters. This will identify events that are thermally reversible.

Most DSC instruments purge the air space around the sample and reference pans with a gas (typically nitrogen or helium). The purge gas serves many functions. Firstly, if the instrument is operated at sub-ambient temperatures, the gas, being dry, prevents condensation or freezing of water. It serves as a heat-transfer medium to ensure the pan and contents are at a temperature as close as possible to the heating block (if greater heat transfer is needed, helium is used as the purge gas). Finally, if any gaseous degradation products are emitted from the pan, the purge gas ensures they are carried out to waste and do not condense on the instrument.

Instruments should be calibrated for temperature and enthalpy upon installation and then performance verified on a daily or weekly basis (recalibration being required if the veriacation test indicates it). Calibration is performed with a certiped reference material (CRM). Typically, CRMs for DSC are highly pure materials with well-established melting points and heats of fusion. IUPAC recommends a number of CRMs for DSC calibration (Table 1). Of these, indium is the most widely used material, although calibration with at least two CRMs, possessing melting points over the range at which measurements are to be performed, is advised. CRMs should be used under an inert gas and disposed of after use (the exception is indium, which can be reused if not heated above 180 $^{\circ}$ C). Gallium will react with aluminium and indium will alloy with gold and so care is needed when selecting pan material. As a general point, uncertainty of heat of fusion measurements will be determined by the uncertainty of the balance used to weigh the sample material.

MTDSC requires selection of additional experimental parameters (underlying heating rate and frequency and amplitude of oscillation). Proper selection is vital to ensure artifacts are not introduced to the data post-deconvolution. In particular, MTDSC assumes that the response of the sample varies linearly

with the modulation in temperature. It also assumes that any changes in the underlying heat ow are slow relative to the time scale of the modulation (this allows averaging of the data, required to recover the underlying heat ow). This means there must be many modulations over the course of a transition (a minimum of six is usually recommended). If these conditions cannot be met then deconvolution cannot be achieved. Melting of a pure material is an example where deconvolution usually fails, because as a material melts its temperature will not rise until melting has θ mished; thus, during melting the temperature of the sample cannot be modulated. Selection of the modulation parameters therefore requires some prior knowledge of the transitions through which the sample will progress and it may be that several experiments will need to be performed with varying parameters to optimise the data. Typical starting values are an underlying heating rate of 2 $^{\circ}\textrm{C min}^{-1}$, frequency 30–60 s and amplitude 0.5–1 \degree C.

Selection criteria for DSC instruments

Table 2 summarises key features of DSC instrumentation and includes criteria to be considered when purchasing DSC equipment. It also provides some guidance on instrumental requirements for different applications, for example measurements above or below ambient temperature.

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photometric analysis

