

Thermal Analysis

TGA / DTA

Outline



- Definitions
- What is thermal analysis?
- Instrumentation & origin of the TGA-DTA signal.
- TGA } Basics and applications
- DTA }
- Phase diagrams & Thermal analysis
 - Thermal analysis, an experimental method to determine phase diagrams.



Nomenclature of Thermal Analysis

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Definition of the field of Thermal Analysis (TA)

Thermal Analysis (TA) is a group of techniques that study the *properties of materials as they change with temperature*

Thermal analysis

- In practice thermal analysis gives properties like; enthalpy, thermal capacity, mass changes and the coefficient of heat expansion.
- Solid state chemistry uses thermal analysis for studying reactions in the solid state, thermal degradation reactions, phase transitions and phase diagrams.



Thermal analysis

... Includes several different methods. These are distinguished from one another by the property which is measured.

- Thermogravimetric analysis (TGA): mass
- Differential thermal analysis (DTA): temperature difference
- Differential scanning calorimetry (DSC): heat difference
- Pressurized TGA (PTGA): mass changes as function of pressure.
- Thermo mechanical analysis (TMA): deformations and dimension
- Dilatometry (DIL): volume
- Evolved gas analysis (EGA): gaseous decomposition products

Often different properties may be measured at the same time:

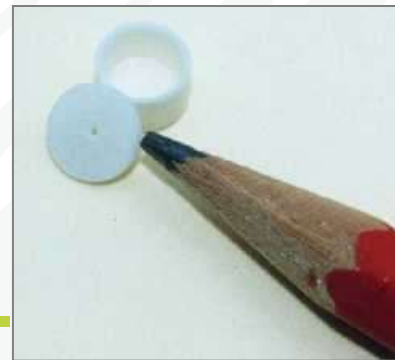
TGA-DTA, TGA-EGA

Instrumentation
&
origin of the TGA-SDTA signal

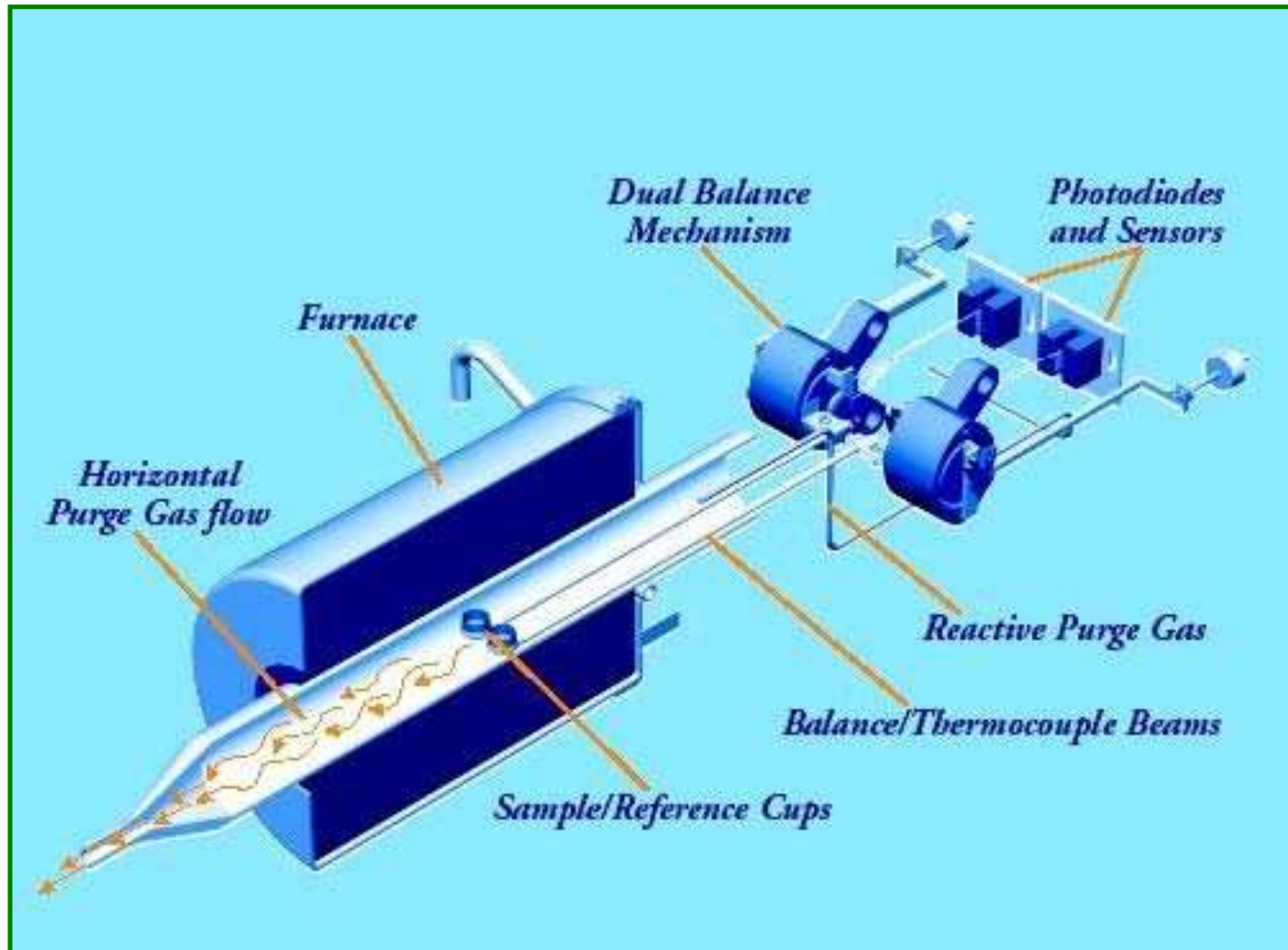


TGA - SDTA

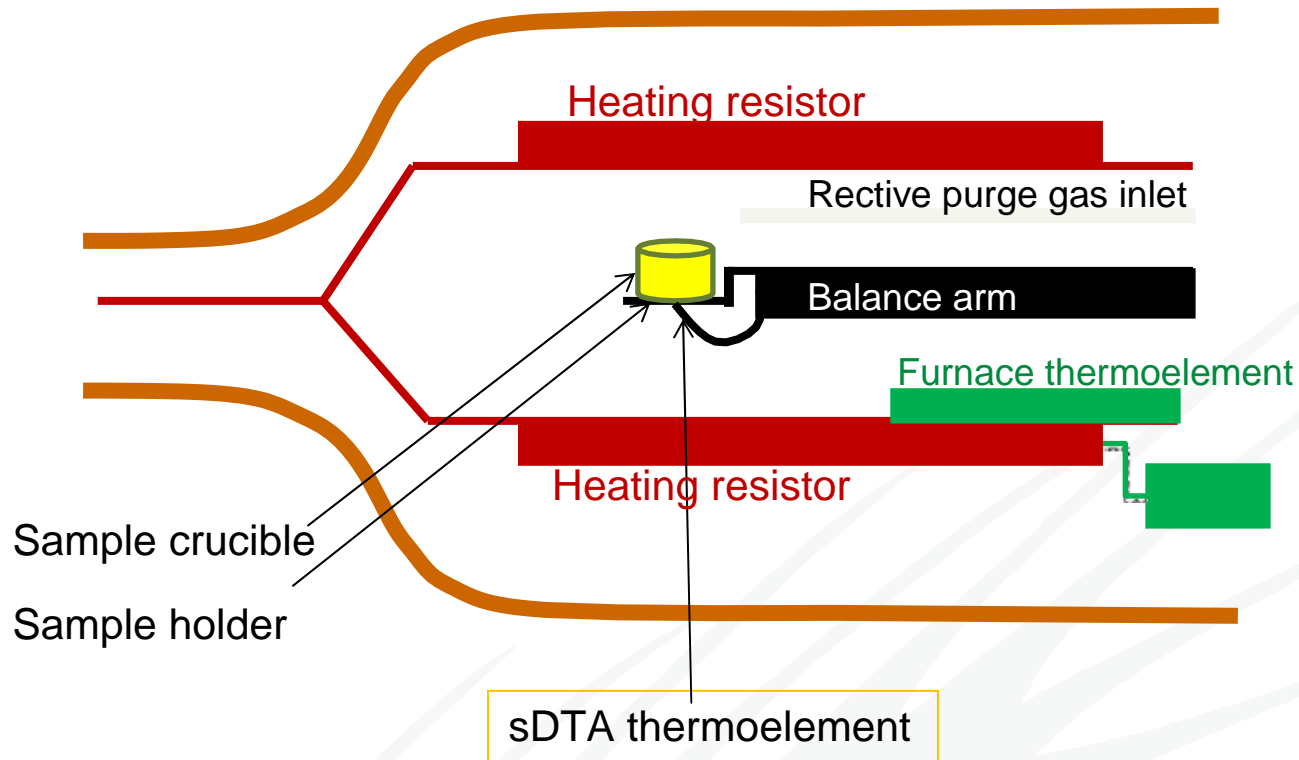
Mettler - Toledo



A modern TGA - DTA



Furnace components

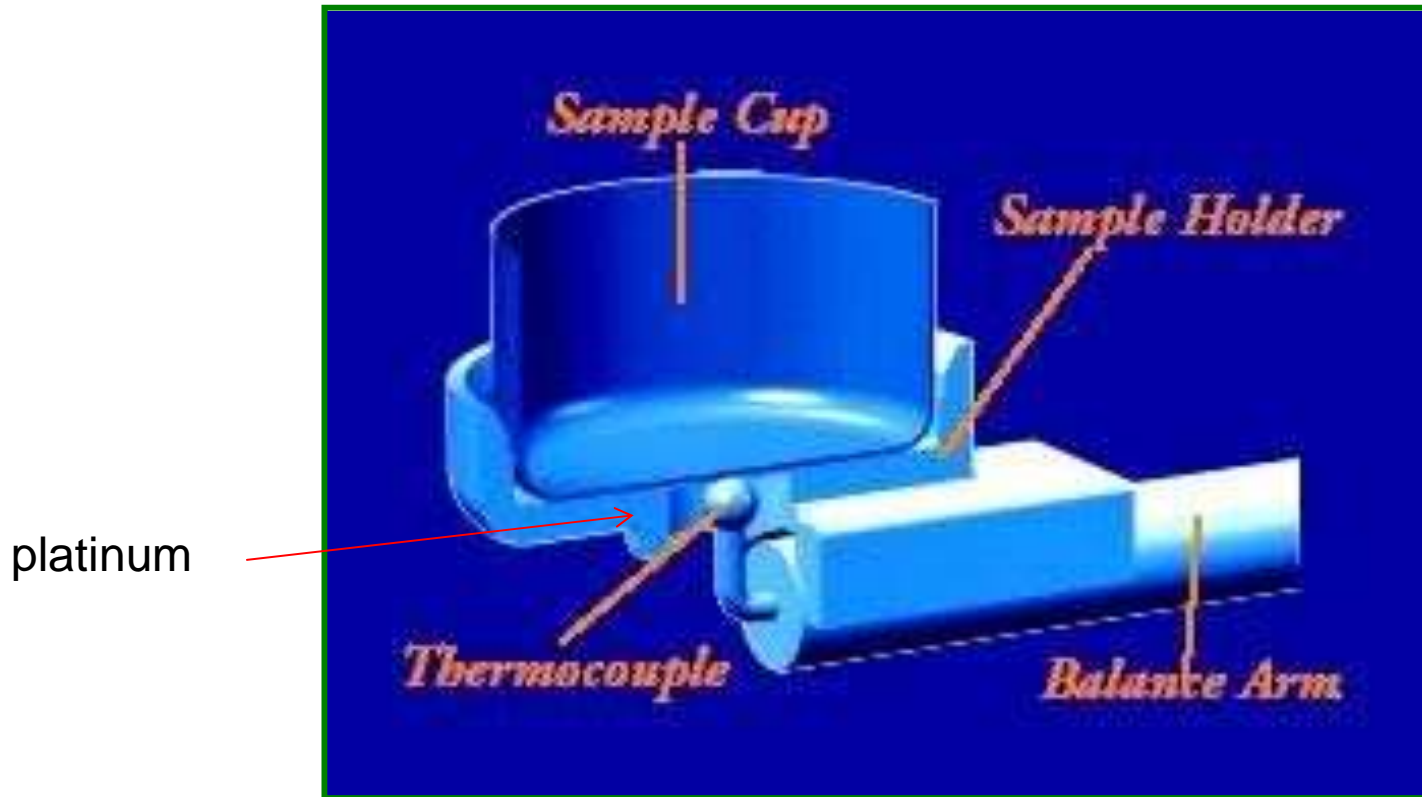


Operating range:
- 200 - 1600 °C

Heating rate:
up to 100 °C/min

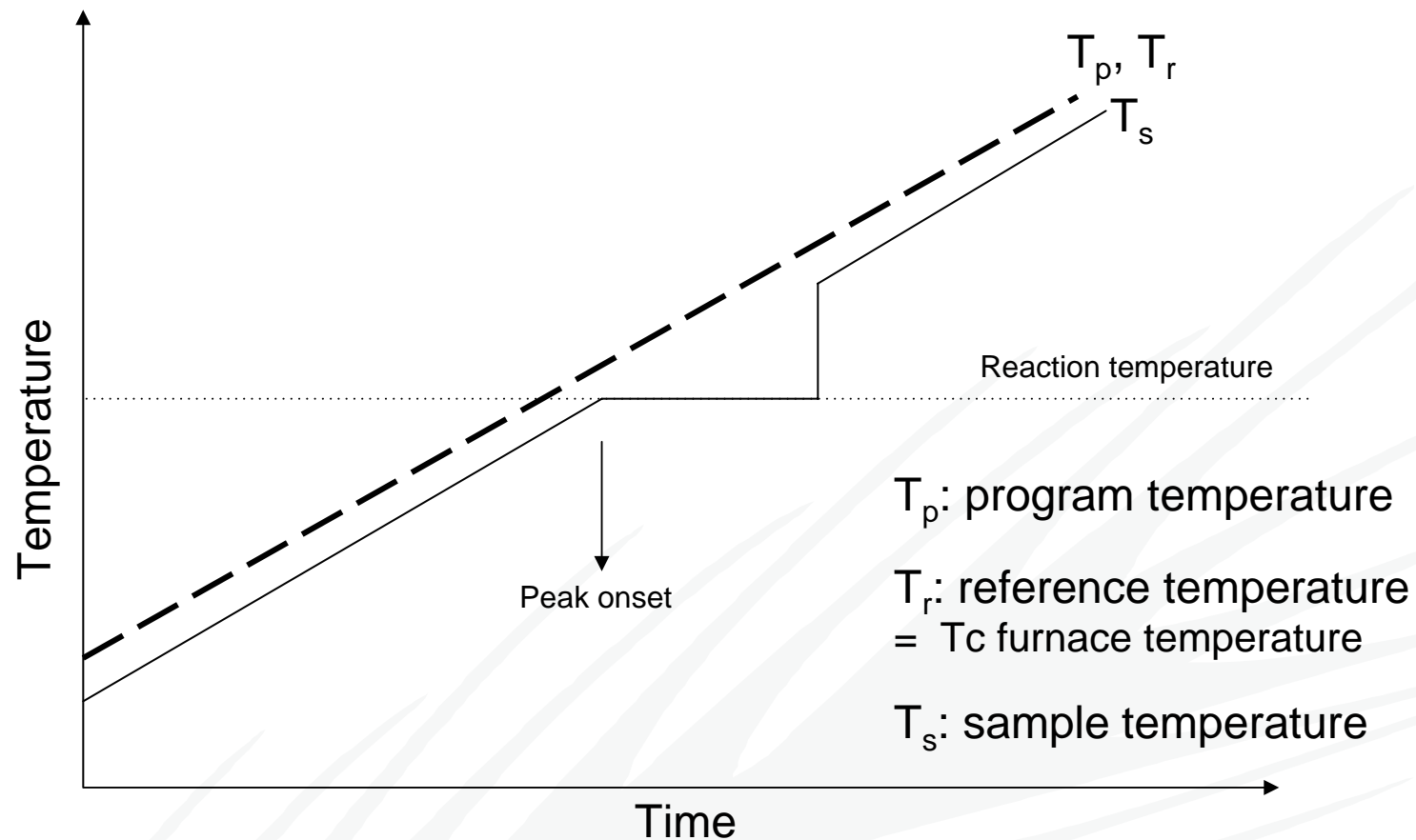
Typical heating rate:
10 – 20 °C/min

Heat transfer from crucible to recording microbalance & thermo elements

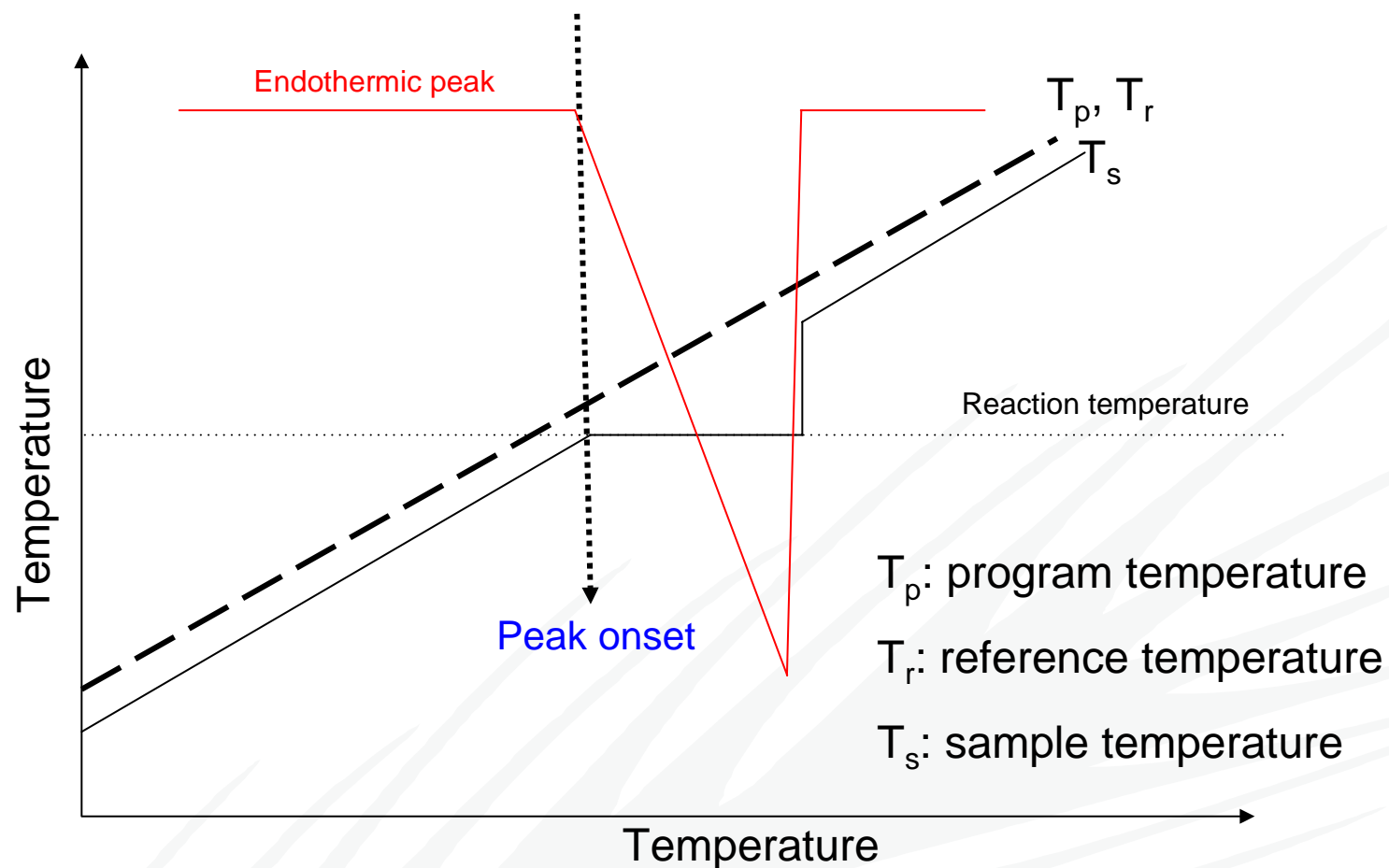


Origin of the TGA-DTA signal

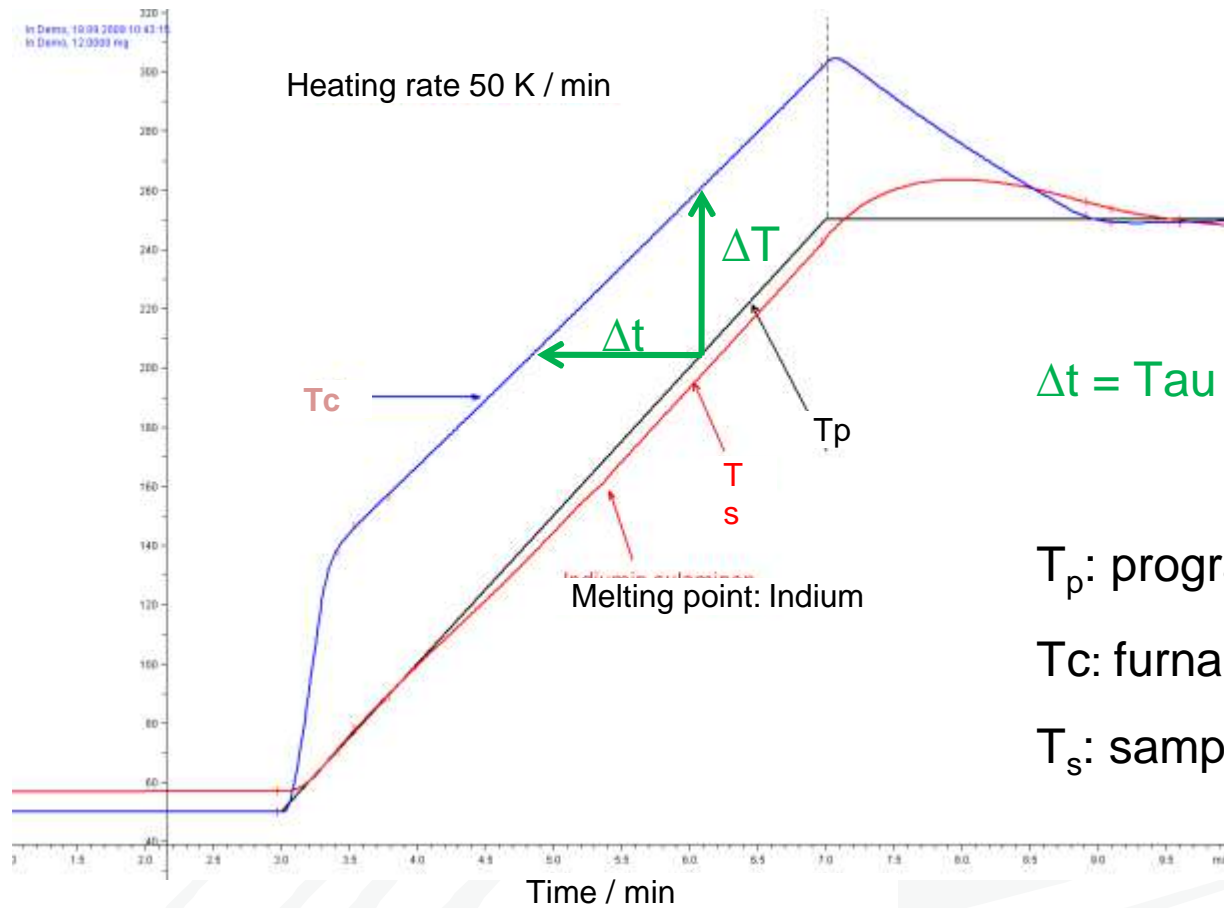
Schematic diagram showing the different temperatures in the DTA during a thermal process.



Origin of the TGA-DTA signal



Origin of the TGA-DTA signal



$\Delta t = \text{Tau lag}$

T_p : program temperature

T_c : furnace temperature

T_s : sample temperature

TGA

Thermo Gravimetric Analysis



TGA, Basics

Measures changes in weight in relation to changes in temperature.

The measured weight loss curve gives information on:

- changes in sample composition
- thermal stability
- kinetic parameters for chemical reactions in the sample

A derivative weight loss curve can be used to tell the point at which weight loss is most apparent



TGA; Phenomena causing mass changes

Physical

Gas adsorption

Gas desorption

Phase transitions

- Vaporization
- Sublimation

Chemical

Decomposition

Break down reactions

Gas reactions

Chemisorption

(adsorption by means of chemical instead of physical forces)

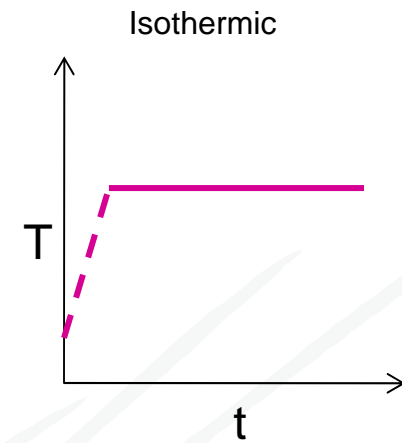
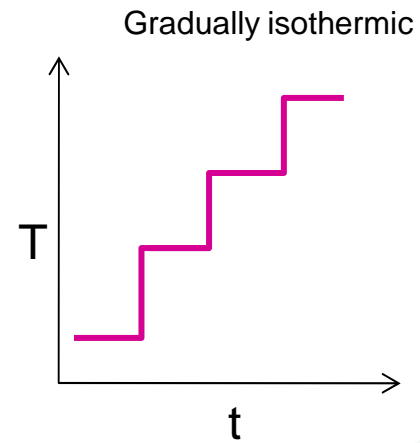
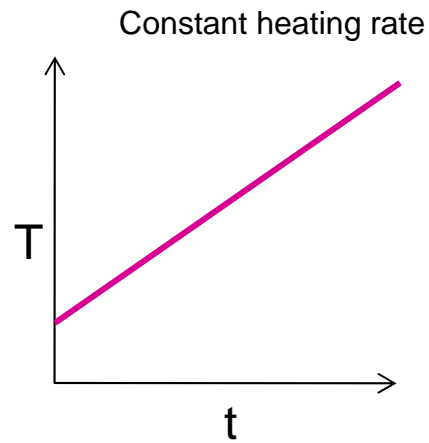


TGA: Applications

- Characterization of
 - Thermal stability
 - Material purity
 - Determination of humidity
- Examination of
 - » Corrosion studies (e.g. oxidation or reactions with reactive gases)
 - » Gasification processes
 - » Kinetic processes



Typical temperature-time programs



Experimental

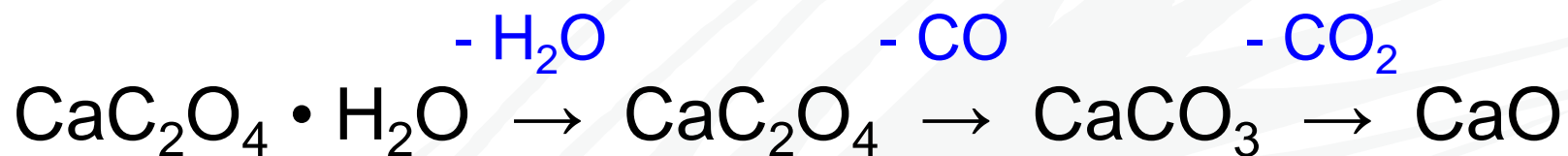
Sample size: 1 – 100 mg (typically 5 – 20 mg)

Heating / cooling rate: 1 – 50 °C / min

TGA

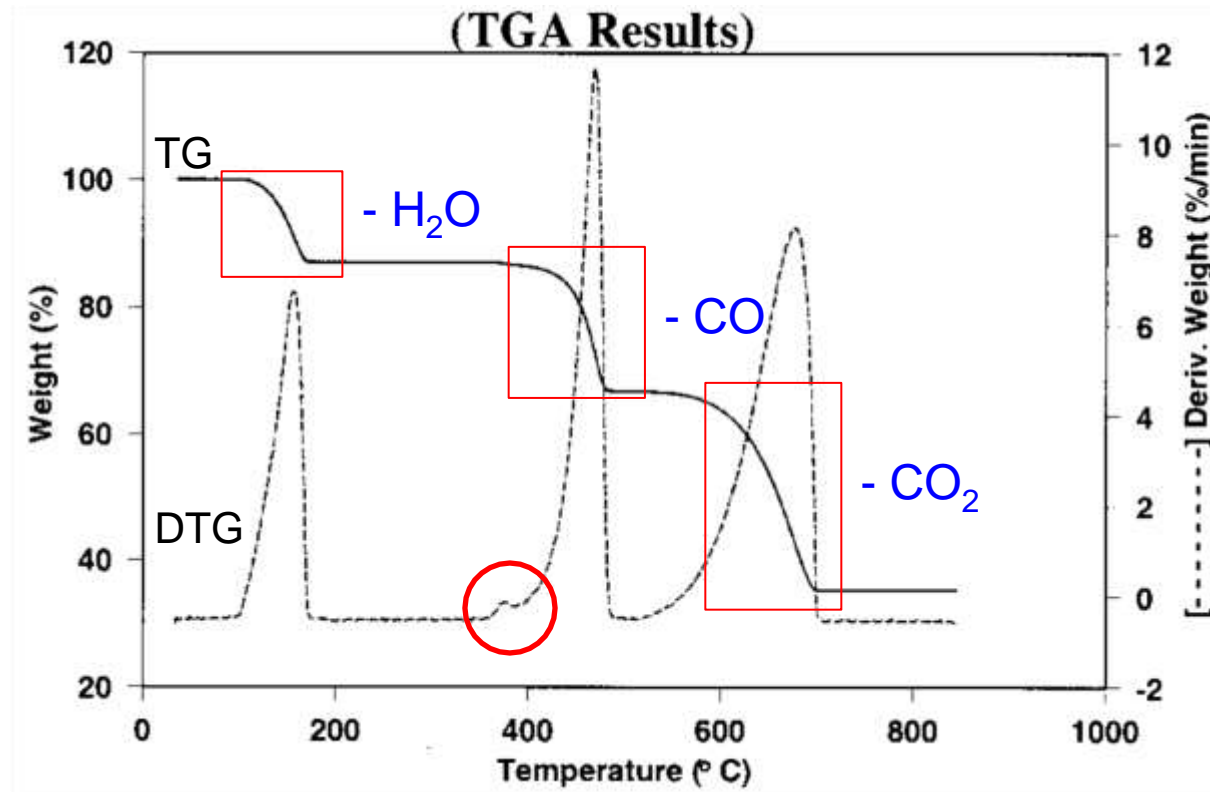
Ex. Decomposition of calcium oxalate monohydrate

- Calcium oxalate monohydrate, a standard material often used to demonstrate TGA performance.
- Exhibits three weight losses with temperature in an inert atmosphere (e.g. N₂).



TGA

Ex. Decomposition of calcium oxalate monohydrate



Absolute confirmation of the decomposition process is possible when the gaseous by products are identified as they evolve, eg. by mass spectrometry (MS).

TGA

Common gaseous components originating from inorganic materials that decompose before the melting point:

H_2O , CO , CO_2 , SO_x , NO_x , Cl_2 , F_2 , CH_3OH , etc.

Also some chemical reactions in solid phase result in gaseous weight loss ex.



Factors affecting the TG curve

Heating rate
Sample size



Increases the temperature at which sample decomposition occurs.

Particle size of sample
Packing
Crucible shape
Gas flow rate

- Affects the progress of the reaction



DTA

Differential Thermal Analysis



DTA, Basics

The material under study and an inert reference are made to undergo identical thermal cycles.



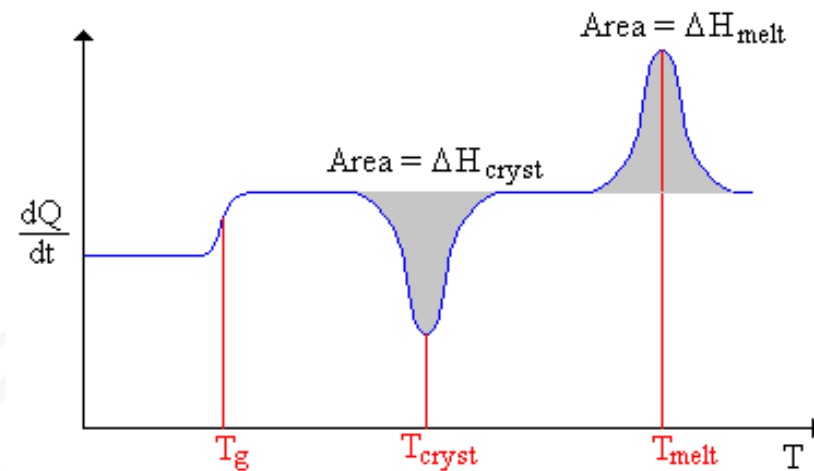
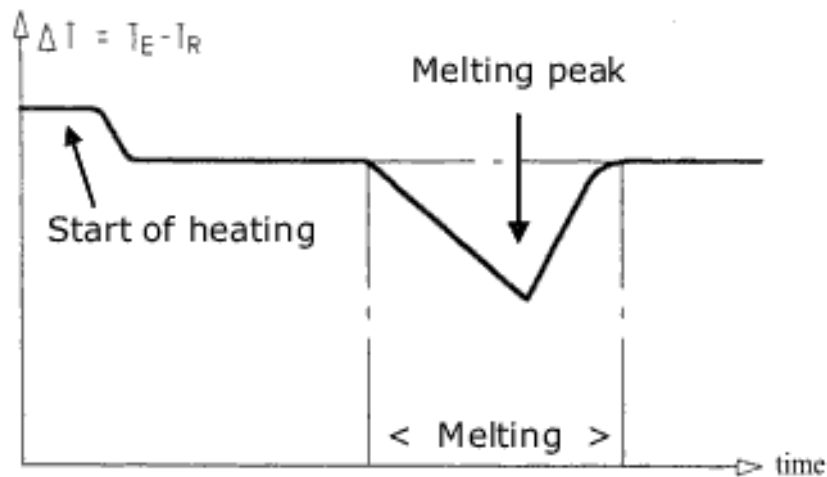
Any temperature difference between sample and reference is recorded.

In this technique the heat flow to the sample and reference remain the same rather than the temperature.



DTA, Basics

➔ The differential temperature is then plotted against time, or against temperature (DTA curve or thermogram).



Crystallization	exothermic	}
Melting	endothermic	

Peak orientation $\uparrow\downarrow$ in DTA thermogram depends on Instrument manufacturer

DTA; Phenomena causing changes in heat / temperature

Physical

Adsorption (exothermic)

Desorption (endothermic)

A change in crystal structure
(endo – or exothermic)

Crystallization (exothermic)

Melting (endothermic)

Vaporization (endothermic)

Sublimation (endothermic)

Chemical

Oxidation (exothermic)

Reduction (endothermic)

Break down reactions
(endo – or exothermic)

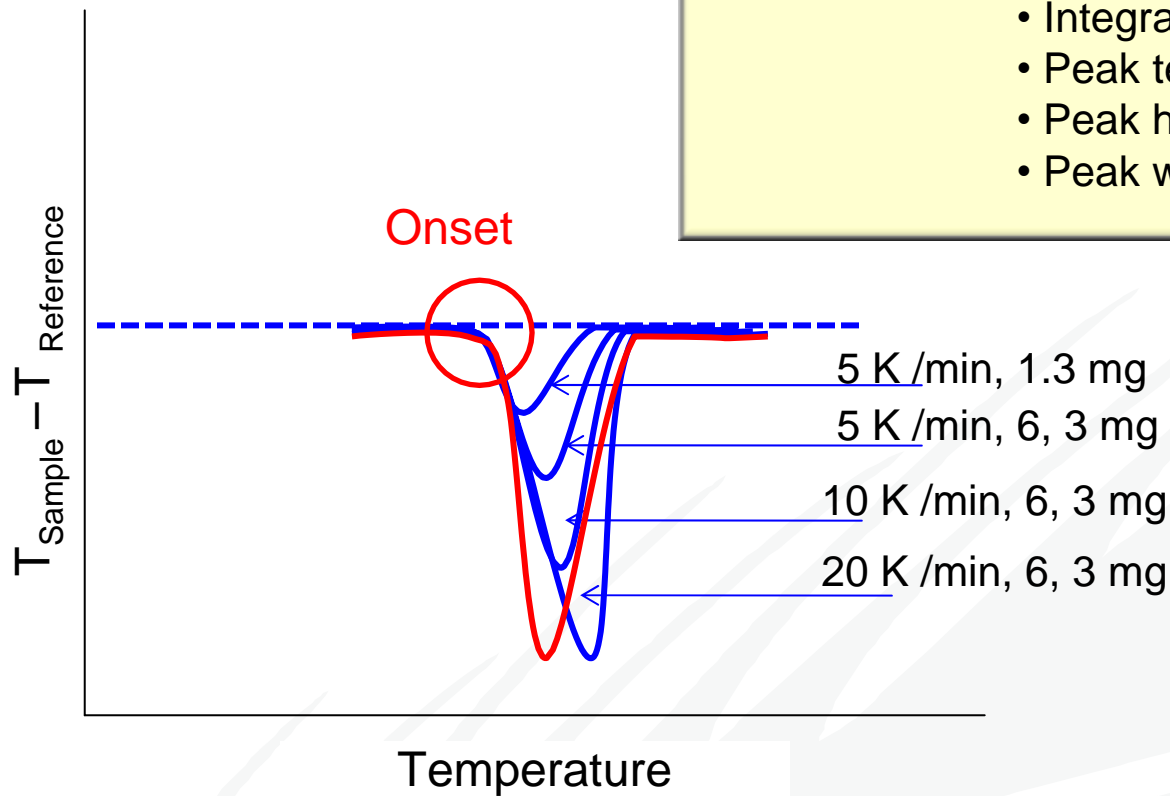
Chemisorption (exothermic)

Solid state reactions
(endo – or exothermic)

Evaluation and interpretation of DTA curves

Typical data obtained from DTA peak evaluation

- Onset - melting
- Endset
- Integral - enthalpy Δh
- Peak temp - melting
- Peak height
- Peak width



Peak temperature is affected by heating rate & sample mass, but not by Δh (enthalpy) and T onset.

TGA- DTA

Keys for successful experimental practice

- Raw materials should be of high purity.
- Fine -grained powder should be used to achieve greater contact area and better equilibrium conditions.
- The time at any temperature must be sufficiently long in order to permit completeness of reactions.



TGA- DTA

Keys for successful experimental practice

Factors affecting the heat transfer, Tau lag & signaling

Crucible

Material

Mass

Volume

Heat capacity

Sample

Mass

Heat capacity

Heat conductivity

Atmosphere



Phase diagrams & Thermal analysis



Phase Diagram

- A phase diagram show conditions at which thermodynamically distinct phases can occur at equilibrium.
- It is determined experimentally by recording cooling rates over a range of compositions.
- Phase transitions occur along lines of equilibrium (=phase boundaries).
 - Solidus = Temp. **below** which the substance is stable in the solid state.
 - Liquidus = Temp. **above** which the substance is stable in a liquid state.

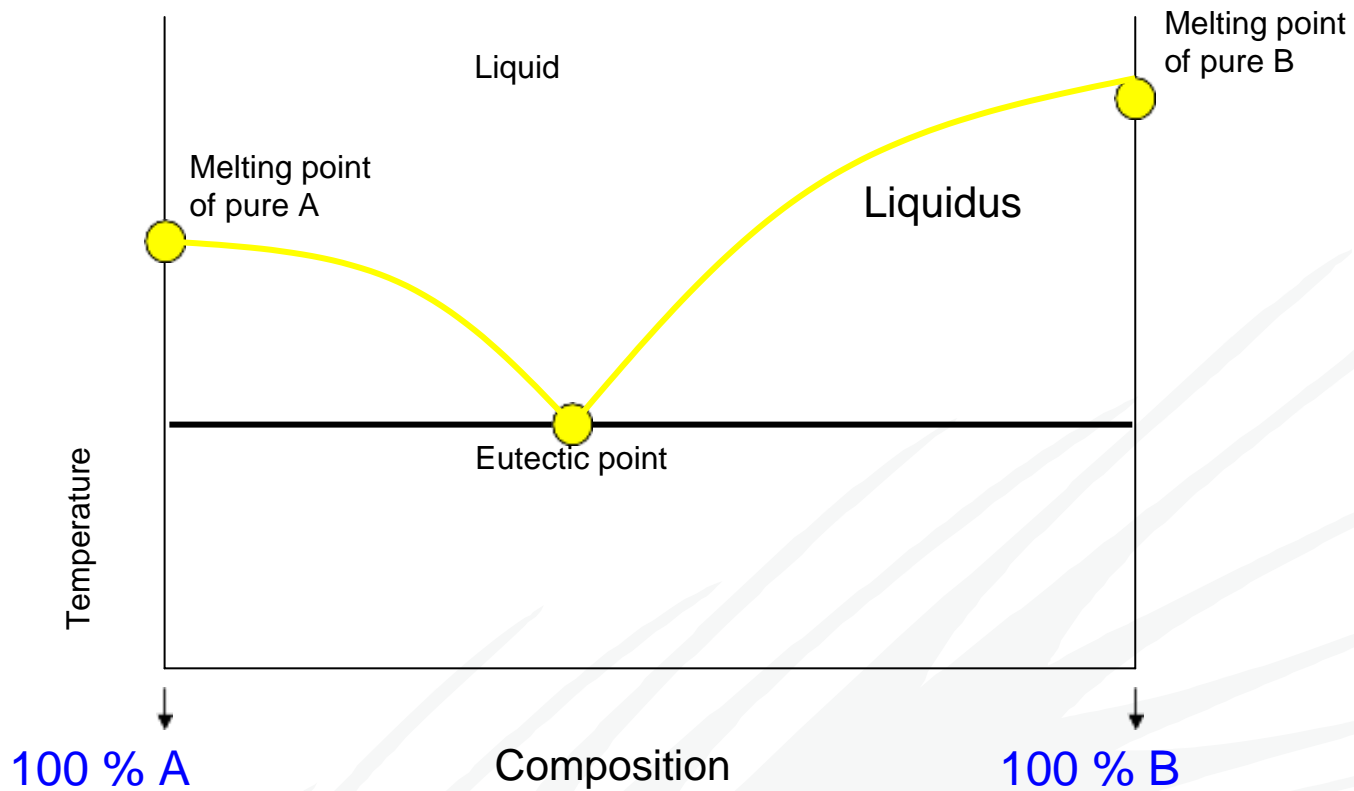


Experimental methods for determining phase diagrams

- Thermal analysis
- High temperature microscopy
- High temperature X-ray diffraction
- Measurement of electrical conductivity as function of temperature.
 - Salt mixtures: solid salts have low conductivity, melts have high.

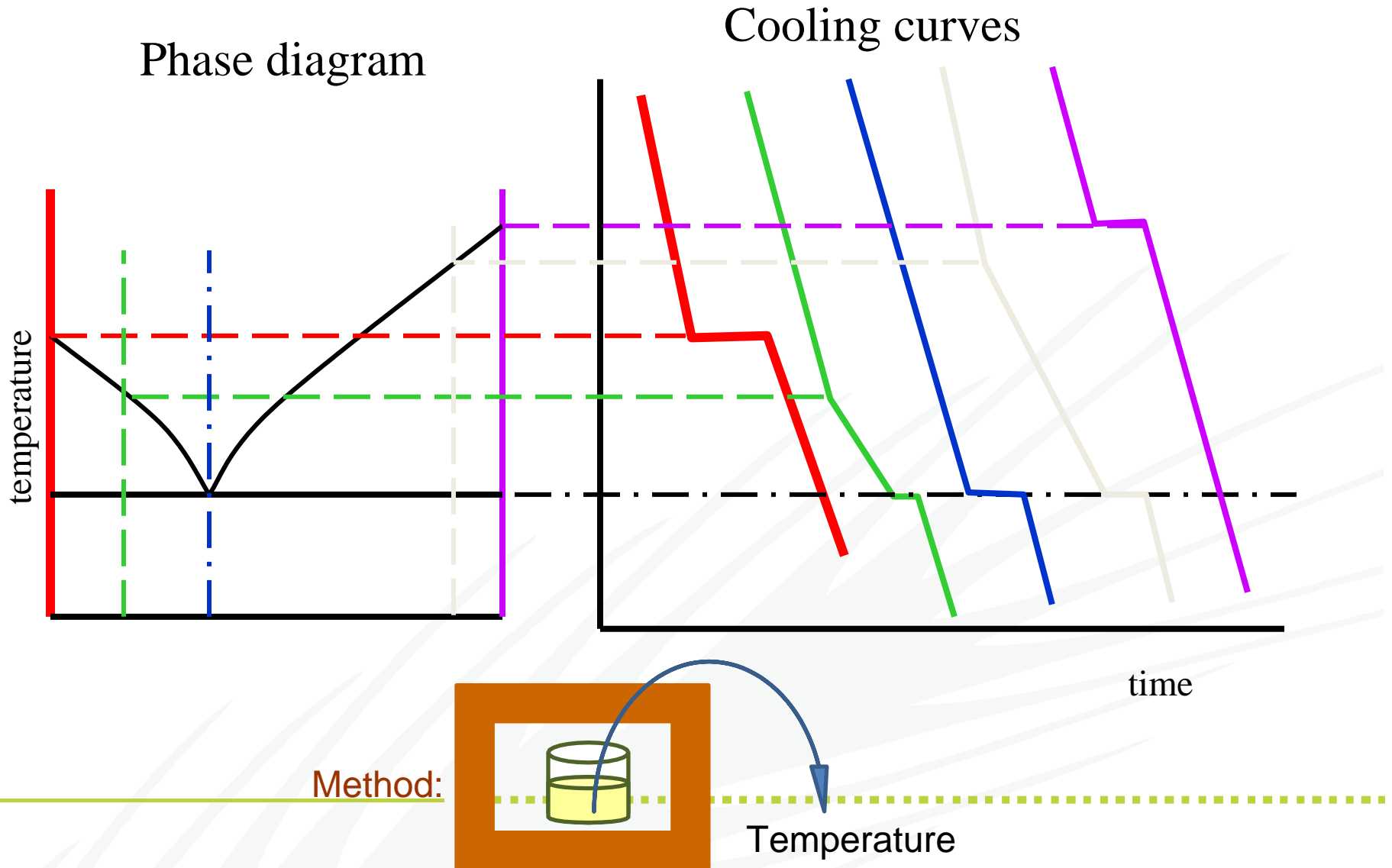


How to build a phase diagram

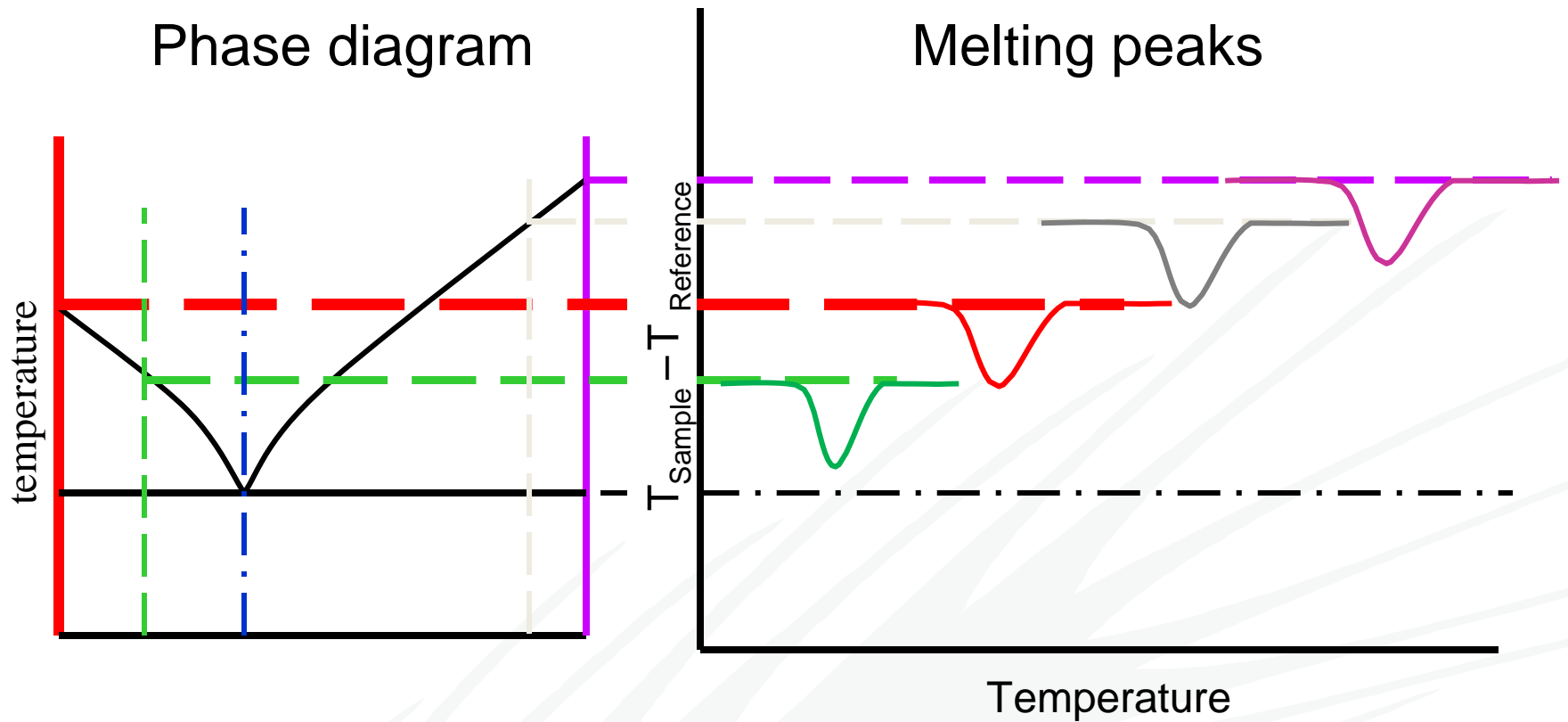


Constructing phase diagrams by experimental methods:

A) from cooling curves



Constructing phase diagrams by experimental methods:
B) from DTA curves



Summary

- Thermal analysis gives information about changes in material properties as function of temperature.
 - Several different TA methods exist; focus on TGA - DTA
 - Combining the two techniques (TGA-DTA) - comprehensive study of a materials thermal behaviour.
 - ▶ While TG only measures changes caused by mass loss, DTA also register changes in material where no mass loss occur, e.g. *crystal structure changes, melting, glass transistion, etc.*
 - Carefullenes required with performance of the experimental procedure to obtain correct weight loss curves and thermograms (e.g. sample preparation, choice of crucible, choice of thermal program)
 - Origin of TG-DTA signal good to know for better understanding of measured data.
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