Anonymous theory: 1761. J. Black: Observed same temperature for ice & water & conjectured about latent" heat.

1797 J. Watt: converled heat -> work as Steam engine

1797 B. Thompson, C. Rumford: work > heat

Calorie fluid thuory: 1783 A. Lavoisier: conjectured heat as invisible fluid.

1824 S. Carnot; Carnot engine > Thermochemistry.

Kinetic throng:

1738 D. Bernoulli: molecular thony of fluid.

1847 J.P. Joule, J. von Mayer, H. von Helmholtz: heat I work equivalent form of energy.

1850 R. Clausius: (a) showed that "caloric fluid theory" in reconciled in "kinetic theory".

(b) First law of Thermodynamics, (c) defined

"entropy.

1854 Thomson or Lord Kelvin -> Thermodynamics

Experimental gos laws:

1662 Boyle's law: Pav

1802 Charle's low; VXT

1802 Gay-Lussac's law:

International scale of temperature

Lord Kelvin introduced lowest temperature independent of property of substance Sulphur point 444.6°C absolute zero - 273°C or OK Antimory point 630.5°C ice point o°C or 273 k Silver point 960.8°C steam point 100°C or 373 k Gold point 1063.8°C

Human body temperature 98.4°F (36,9°C (remember, least count is oic) relation between C, F, R (Reumer Scale)

$$\frac{C}{5} = \frac{F-32}{9} = \frac{R}{4}.$$

Basics of Thermometry

- liquid Thermometer: Hg in glan, 6's max-min, Beckmann, Index
- Gas Thermometer: Callendar's Air, Hydrogen
- Metal Thermometer: Platinum resistance,
- d) Thermoeletric Thermometer: thermocouple generates C.m.f. e) Radiation Thermometer: e = at f bt2

principle of thermometry: let A = thermometric property

For uniform variation of A with temperature T AT = Ao (1+ dT), Ao = value of A at Oc A100 = value of A at 100C.

A100 = A0(1+ 100x)

$$T = \log \frac{A_{T} - A_{0}}{A_{100} - A_{0}}$$

What is A? If you take cimple Hg-in Slam thermometer then A = length of Hg column. For Pt-resistance thermometer A = resistance R of Pt. For constant volume gas thermomder A = pressure of air.

Hg in glass thermometer

pros (a) liquid range [-38.87°c, 356°c]

- Low thermal capacity & Ligh conductivity
- uniform expansion, d) shining liquid (easy to see).

(a) high specific gravily, (less sensitive)

(b) high surface tension (molion is jerkey)

unsuitable for low T.

pros (a) liquid range [-112°c, 78°c]

- (high coefficient of expansion (more sensitive)
- (c) low surface tension (motion is smooth)

cons unsuitable for high temperature.

Gas thermometer (a) Very high coefficient of expansion (more sensitive

- (b) low thermal capacity, low specific gravily
- (c) constant volume / constant pressure the mometer.

Platinum Resistance: Resistance smooth in range $[-200^{\circ}c]$ thermometer least count = $0.01^{\circ}c$ $R_t = R_0(1+dT)$

low temperature Thermometry

o'c to 630c -> all

-20c to 0'c -> liquid, gas,

resistance, thermosuple,

vapour pressure.

-272'e to-20'c -> resistance, vapour

High temperature thermometry

630°c to 1000°c → liquid, gas,
resistance, thermocouple.

1600°c to 1600°c → gas, thermocouple.

1600°c to 3000°c → thermocouple.

pyrometer.

below - 272°C - magnetic

1. At what temperature do the Fahrenheit & celsium scale give the same reading?

2. In figure beside, which line represents $\frac{75}{5} = \frac{5-32}{9}$?

pressure, magnetic

3. In a constant volume gos thermomeler, pressure of air at o'c is 80 cen f at 100 c is 109.3 cm. Calculate the temperature of a hot bath, inwhich when the thermometer is immersed shows a pressure 100 cm.

Colorimetry Definition of Calorie (heat) & quantily of heat needed to raise temperature of 1 gm of water by 1°c. But heat 5°c > 6°c ≠ 80°c > 81°c.

Mean calorie heat [oc -> 100c]/100.

15° Calorie heat 14.5° -> 15.5°

B. Th. U. quantily of heat needed to raise temperature of 116 of water by 1°F. 1 B. Th. U. = 252 cals

1. C. H. U. = 453.6 cals. (centrigrad heat unit)

H = M S T = CT

heat mass specific thermal of substance heat capacity

Basic principle of Calorimetry conservation of head energy or sum total of (body + head bath) is conserved.

Specific heat of solid: Method of mixture, copper block colorimeter, Electrical method (Lindemann colorimeter), Latent Locat (Bunsen Fee colorimeter).

Specific heat of liquid : Method of mixture, Bursen ice calorimeter, Method of cooling,

Nernst calorimeter, continuous flow calorimeter.

Specific heat of gas: # copy constant pressure or volume?

C = d8 -> suppose system expands -> temperature falls
to prevent, d8 heat added so dT=0.
C = d8 -> &

→ suppose system compresses -> temperature rise dT without the need of dg :: dg=0 C= = 0. cp = $\left(\frac{dg}{dT}\right)_{P}$, $C_{V} = \left(\frac{dg}{dT}\right)_{V}$.

definition: at P = constant, increase temperature of 1 gm of gas through 1°c. If 1 gm molecule of gas is taken \Rightarrow molar specific heat. $C_{P} = MC_{P}$ $C_{P} > C_{V}$? # heat transfer at V = constant, dV = 0. dW = PdV = 0no work done by the gas. $C_{P} - C_{V} = \frac{R}{J}$ So heat supplied transferred to thermal energy that the paint # heat transfer at P = constant, $dW = PdV \neq 0$. $C_{P} - C_{V} = R$ so heat supplied \Rightarrow thermal energy + external work sothermal: T = constant

Isothermal: T = constant [PV=RT = constant]

transformation

Adiabatic : Q = constant, $dQ = C_V dT + P dV = 0$ transformation

Now from PV=RT, P dV + V dP = R dT = (CP - CV) dTput dT in O: $dQ = C_V \frac{P dV + V dP}{CP - CV} + P dV = 0$ (as Q = condowb) $C_V V dP + CP P dV = 0$

Integrating both sides, $\sqrt{lnV} = -\frac{dP}{P}$, (put $\sqrt{s} = \frac{CP}{CV}$) $\sqrt{lnV} = -lnP + lnC$. $\sqrt{PV} = constant$

Corollary: PT relation: put PV=RT, $P(\frac{PT}{P})^2 = C \Rightarrow P^{1-7}T^2 = corollard$ VT relation: put PV=RT, $\frac{PT}{V}V^2 = C \Rightarrow V^{3-7}T = corollard$

Suppose dry air is enclosed in a cylinder filled with piston.

Initial temperature of air is 30°C. Find the change in

temperature if the gas is compressed (a) slowly, (b) suddenly

to one-half of its volume? What'll be the pressure?

(a) piston moves inwards -> gas compresses



no change < heat goes to < temperature rices in surrounding by temperature conducting wall (isothermal) slowly T=30c

(b) Temperature rise rate >>> heat diffusion rate to surrounding adiabatie process - A no & transfer.

So $P_1V_1' = P_2V_2''$ & given $V_2 = \frac{V_1}{2}$ $\therefore P_2 = 2^{\gamma}P_1 = 2^{1-\gamma}P_1 = 2.6P_1 = \frac{2.6 \text{ otn}}{2.6P_1} \text{ when } P_1 = 1 \text{ atm.}$ Compare with isothermal, $P_2 = 2P_1 = \frac{2 \text{ atm.}}{2}$ Again from TV relation, $T_1V_1'' = \overline{T_2V_2'}$

 $T_2 = T_1 \left(\frac{V_1}{V_2} \right)^{2/-1} = 2^{0.4} T_1 = 1.319 \times (273 + 30) \times$ $= 399.7 \times = (399.7 - 273)^{\circ} = 127^{\circ} = 1.27^{\circ} =$

- 4. Air in a Wisson's cloud chamber at 20°c is abruptly expanded to 1.4 times its initial volume. Calculate the final temperature. Given $\beta = 1.4$.
 - 5. At 10° dynes/cm² pressure, a gas (=1.4) expands isothermally until its volume is double of the initial volume. Then it adiabatically expands until its volume is redoubled. Calculate the final pressure of the gas.