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 law; V & T

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 Antimony 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 (semember, 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+ 100%)

$$T = \log \frac{A_{\tau} - A_0}{A_{100} - A_0}$$

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]

6) Now 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 thermometer.

Platinum Resistance: Resistance smobth in range  $[-200^{\circ}c]$  thermometer least count  $= 0.01^{\circ}c$   $R_{t} = R_{0}(1+\Delta T)$ 

Low temperature Thermometry

octo 630c -> all

-20c lo oc -> liquid, gas,

resistance, thermocouple,

vapour pressure.

-272 e to-20 e -> resistance, vapour pressure, magnetic

below - 272° - magnetic

High temperature thermometry

630c to 1000c - liquid, gas,

resistance, thermocouple.

1000c to 1600c + gas, thermocouple.

1600°c to 3000°c -o thermocouple, pyrometer.

HW 1. At what temperature do the fahrenheit & celsium scale of give the same reading?

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

3. In a constant volume gors thermometer, 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) D quantity 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 colorie 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. CoHoUo = 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 heat energy or sum total of (body + heat bath) is conserved.

Specific heat of solid: Method of mixture, lopped block calorimeter, Electrical method (Lindemann calorimeter), latent hout (Bunsen Fee calorimeter).

Specific heat of liquid & Method of mixture, Bursen ice colorimeter, Method of cooling, Nernst Calorimeter, continuous flow calorimeter.

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

C = d8 -> suppose system expands -> temperature falls
toprevent, dg heat added so dT=0.
C = d8 -> 00

→ suppose system compresses -> temperature rise dT without the need of dQ :: dQ = 0 C = 0 = 0. cp =  $(\frac{dg}{dT})_p$ ,  $C_V = (\frac{dg}{dT})_V$ .

definition: at P = constant, increase temperature of 1 gm of gas through 1°c. If 1 gm molecule of gas 5 taken  $\Rightarrow$  molar specific heat.  $C_p = MC_p$   $C_p > C_V ?$  # heat bransfer at V = constant, dV = 0. dW = PdV = 0no work done by the gas.  $C_p - C_V = \frac{P}{J}$  So heat supplied transformed to thermal energy.

(heat) # heat transfer at P = constant,  $dW = PdV \neq 0$ .  $C_p - C_V = R$ So heat supplied  $\Rightarrow$  thermal energy + external work.

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

transformation  $Adiabatie : Q = constant, dQ = C_V dT + P dV - D$  transformation  $Now \text{ from } PV = RT, P dV + V dP = R dT = (C_P - C_V) dT$   $Put dT \text{ in } D : dQ = C_V \frac{P dV + V dP}{C_P - C_V} + P dV = 0 \text{ (as } Q = condant)$ 

 $C_{V}VdP + C_{P}PdV = 0$   $C_{V}VdP + C_{P}PdV = 0$   $C_{V}VdP + C_{P}PdV = 0$   $C_{V}VdP + C_{P}PdV = 0$ 

Integrating both sides, & lov = -lnp+ lnc.

PV = constant

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

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  $P_1V_1' = P_2V_2'' + given V_2 = \frac{V_1}{2}$   $\therefore P_2 = 2^{7}P_1 = 2^{104}P_1 = 2.6P_1 = \frac{2.60 \text{ atm}}{2.60 \text{ when } P_1 = 1 \text{ atm}}$ . Compare with isothermal,  $P_2 = 2P_1 = \frac{2}{2} \text{ atm}$ .

Again from TV relation,  $T_1V_1^{V-1} = T_2V_2^{V-1}$   $T_2 = T_1\left(\frac{V_1}{V_2}\right)^{V-1} = 2^{0.4}T_1 = 1.319 \times (273+30) \times 10^{-1}$   $T_1 = 1.319 \times (273+30) \times 10^{-1}$  $T_2 = T_1\left(\frac{V_1}{V_2}\right)^{V-1} = 2^{0.4}T_1 = 1.319 \times (273+30) \times 10^{-1}$ 

- 4. Air in a Wisson's cloud chamber at 20°C is absuptly 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.