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## Chemical Kinetics

- It is branch of chemistry which deals with rate of chemical reaction and the factors affecting rate of chemical reaction like temp., conc., pressure, catalyst, etc.

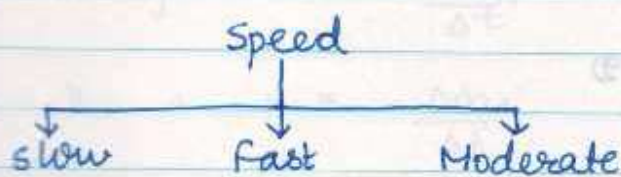


- ⇒ spontaneity / feasibility is predicted by Thermodynamics
- ⇒ extent of Rxn is predicted by chemical eq.
- ⇒ Rate of chemical Rxn is determined by chemical kinetics.

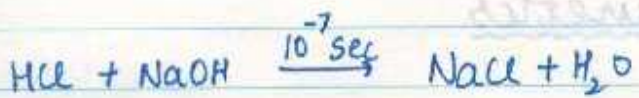
### \* Classification of chemical Rxn :-

Rxn can be classified on the basis of

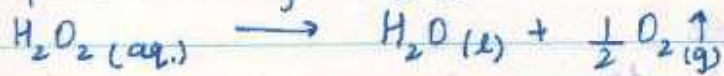
- Speed
- Mechanism



- Slow Rxn : These are the rxns which take place at a very low rate. For eg, rusting of iron.
- Fast Rxn : These are the rxns which occur instantaneously within a fraction of time. For eg, acid-base rxn.



- Moderate Rxn: These are the reactions which are neither very slow nor instantaneous. For eg, decomposition of  $\text{H}_2\text{O}_2$ .

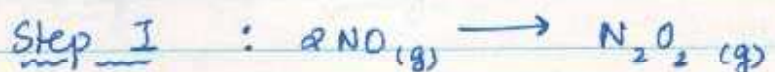
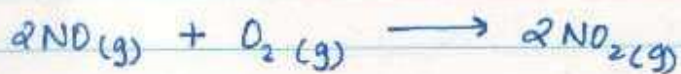


### Mechanism

$\downarrow$   
Simple  
 $\downarrow$   
Elementary

$\downarrow$   
Complex

- Elementary Rxn: These are the reactions which get completed in single step.
- Complex Rxn: These are Rxns which take place in more than one step & each step of rxn will be an elementary rxn. For eg,



- Note:- Just by seeing rxn, we cannot predict whether it is simple or complex rxn.  
- We generally study moderate rxns.

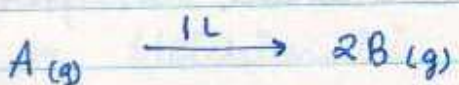
- Rate of Rxn: change in amount of reactant or product per unit time.

- Rate of appearance of Product (ROA):

Amount of product formed per unit time.

- Rate of Disappearance of reactant (ROD):

Amount of reactant disappeared / reacted per unit time.



$t=0$	100 mol	-
$t=1\text{ sec}$	90 mol	20 mol

If  $v$  is constant,

$$\text{ROD of } A = -\frac{\Delta n_A}{\Delta t} = -\left(\frac{n_{t_f} - n_{t_i}}{t_2 - t_1}\right) = 10 \text{ mol/sec}$$

$$\text{ " " " } = -\frac{\Delta m_A}{\Delta t} = -\frac{\Delta [A]}{\Delta t}$$

$$\text{ " " " } = -\frac{dp_A}{dt}$$

$$\text{ROA of } B = \frac{\Delta [B]}{\Delta t} = \frac{20 - 0}{1} = 20 \text{ M/sec}$$

If  $v$  is variable,

$$\text{ROD of } A = -\frac{\Delta n_A}{\Delta t}; \quad -\frac{\Delta m_A}{\Delta t}$$

V → constant



100 -

90 20

$$[A]_i^{\circ} = 10M$$

$$[A]_f = 9M$$

V → Variable



100 - V → 10L

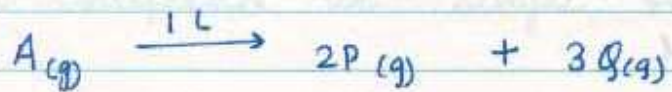
90 20 V → 5L

$$[A]_i^{\circ} = 10M$$

$$[A]_f = 18M$$

Note: If volume is variable then RDA or ROD is expressed in terms of moles or mass but if volume is constant then it can also be expressed in terms of concentration or partial pressure.

e.g.



t=0 100 0 0

t=10s 90 20 30

$$\text{ROD of } A = 1 \text{ mol/s} = 1M/s = -\Delta[A]/\Delta t$$

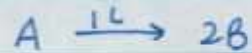
$$\text{RDA of } P = 2 M/s = \Delta[P]/\Delta t$$

$$\text{RDA of } Q = 3 M/s = \Delta[Q]/\Delta t$$

$$\text{Rate of Rxn} = -\frac{\Delta[A]}{\Delta t} = \frac{1}{2} \frac{\Delta[P]}{\Delta t} = \frac{1}{3} \frac{\Delta[Q]}{\Delta t} = 1 M/s$$

In general,

$$\text{ROR} = -\frac{1}{a} \frac{\Delta[A]}{\Delta t} = -\frac{1}{b} \frac{\Delta[B]}{\Delta t} = -\frac{1}{c} \frac{\Delta[C]}{\Delta t} = \frac{1}{d} \frac{\Delta[D]}{\Delta t}$$

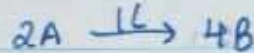


t=0 100

0

t=10 90

20



t=0 100

t=10 90

20

$$\text{ROD of } A = 1 \text{ M/s}$$

$$\text{ROA of } B = 2 \text{ M/s}$$

$$\text{ROR} = 1 \text{ M/s}$$

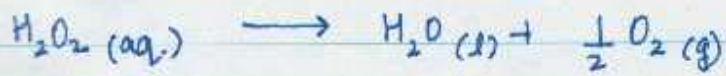
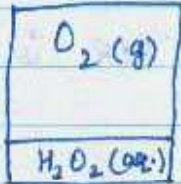
$$\text{ROD of } A = 1 \text{ M/s}$$

$$\text{ROA of } B = 2 \text{ M/s}$$

$$\text{ROR} = \frac{1}{2} \text{ M/s}$$

- ROA or ROD are actual qty. and independent of stoichiometric coeff. however ROR is theoretical qty. & it will depend on stoichiometric coeff.

It means a rxn can have more than one ROR depending on its balancing. So whenever ROR is asked a balanced chemical rxn should be given. or ROR is defined only for balanced chemical eq".



$$\text{ROR} = -\frac{\Delta [\text{H}_2\text{O}_2]}{\Delta t} = \frac{2\Delta[\text{O}_2]}{\Delta t}$$

due to diff. volumes taken into consideration ..

$$\boxed{\text{ROR} = -\frac{\Delta n_{\text{H}_2\text{O}_2}}{\Delta t} = \frac{2n_{\text{O}_2}}{\Delta t}}$$

- Units of ROR :-

$\text{mol s}^{-1}$

$\text{M s}^{-1}$  or  $\text{mol L}^{-1}\text{s}^{-1}$

$\text{atm s}^{-1}$  or  $\text{bar s}^{-1}$

- Conversion :-

$$P_A V = n_A R \theta T$$

$$P_A = \left( \frac{n_A}{V} \right) RT$$

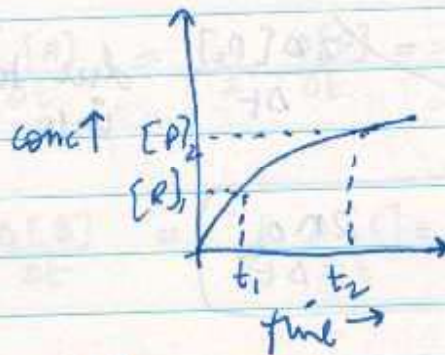
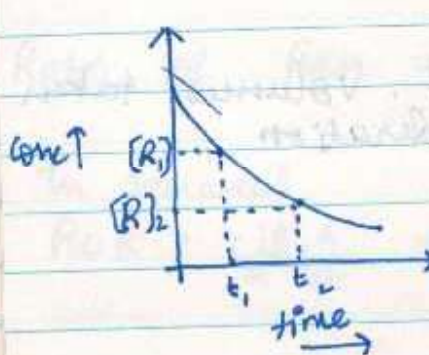
$$\Rightarrow \frac{1}{RT} \times \frac{\Delta P_A}{\Delta t} = \frac{\Delta [A]}{\Delta t}$$

- Types of Rate of Rxn :-

i) Average ROR

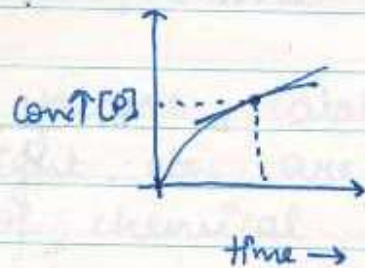
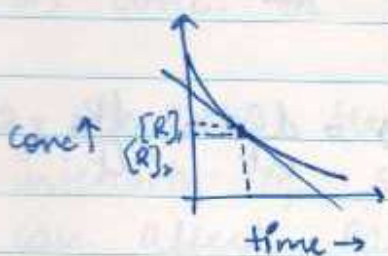
ii) Instantaneous ROR

a) Average Rate of Rxn : If ROR is measured over a large interval of time.



$$\text{R.O.R} = -\frac{\Delta [P]}{\Delta t} = \frac{\Delta [P]}{\Delta t}$$

B) Instantaneous Rate of Reaction: If rate of reaction is measured over a very short interval of time.



$$\text{ROD of } R = -\frac{d[R]}{dt}$$

$$\text{ROA of } P = \frac{d[P]}{dt}$$

$$\text{ROR} = -\frac{d[R]}{dt} = \frac{d[P]}{dt}$$

Q. For the reaction



The no. of moles of B increases by  $6 \times 10^{-3}$  mol in 10 sec in 10 L vessel. Calculate  
 a) ROA of B & C  
 b) ROD of A  
 c) ROR

$$\rightarrow \text{ROA of } B = \frac{6 \times 10^{-3}}{100}$$

$$\text{ROD of } A = \frac{4 \times 10^{-3}}{100}$$

$$C = \frac{8 \times 10^{-3}}{100}$$

$$\text{ROR} = \frac{2 \times 10^{-3}}{100}$$



\* Factors affecting Rate of Rxn :-

- i) Nature of Rxn components
- ii) Temperature
- iii) Conc. or partial pressure

Besides these factors, catalyst, surface area, intensity of sunlight, etc. are other factor which can affect Rate of chemical rxn.

\* Rate law :-



$$\text{Rate of Rxn} \propto a_A^\alpha a_B^\beta$$

$$\boxed{\text{Rate of Rxn} = k a_A^\alpha a_B^\beta} \rightarrow \text{Rate law expression}$$

where,

$a_A, a_B$  → active masses of A & B resp.

$\alpha, \beta$  → experimentally determined factors

$k$  → Rate constant / velocity constant / specific Rate

• Case I : In case of liq. sol<sup>n</sup> active masses are replaced by conc.

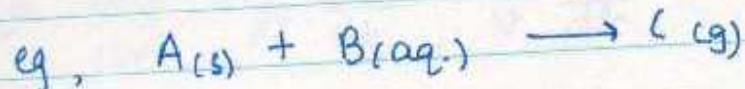
$$\text{Rate} = k [A]^\alpha [B]^\beta$$

Case II : In case of gases active masses are replaced by either conc. or partial pressure

$$\text{Rate} = k P_A^{\alpha} P_B^{\beta}$$

$$\text{Rate} = k' [A]^{\alpha} [B]^{\beta}$$

Case III : In case of pure liquid, pure solid and excess solvent active masses remain constant or replace by unity.

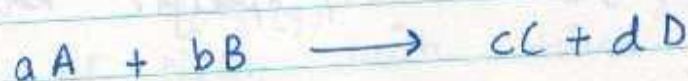


$$\text{Rate} = k a_A^{\alpha} a_B^{\beta}$$

$$= k' a_B^{\beta} = k' [B]^{\beta}$$

\* Order of Rxn :-

as sum of powers of molar conc. or partial pressure of each components in experimentally determined rate law expression.



(iv)  $\boxed{\text{Rate} = k [A]^m \cdot [B]^n} \rightarrow \text{Rate law expression}$

where,

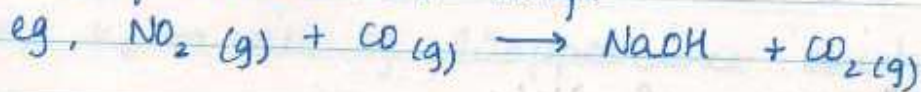
$m \rightarrow$  Order w.r.t. A

$n \rightarrow$  Order w.r.t. B

$m+n \rightarrow$  Order of Rxn

Some imp. points regarding Order of Rxn :-

- It is determined experimentally.
- Order of Rxn can be +ve, -ve, zero or fraction
- Order as well as mechanism of rxn may get changed on changing reaction conditions like pressure or temp.

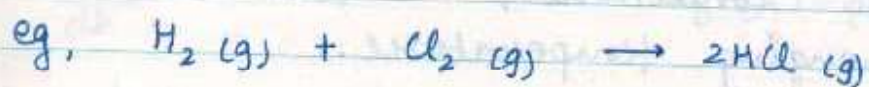


$$\text{Rate} = k [\text{NO}_2][\text{CO}]$$

Above  $500^\circ\text{C}$

$$= k [\text{NO}_2]^2$$

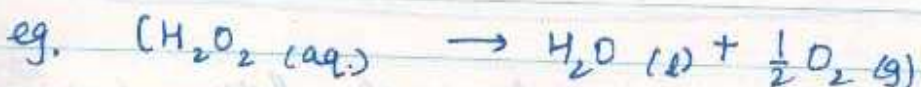
Below  $500^\circ\text{C}$



$$\text{Rate} = k$$

order  $\rightarrow 0$

Rate constant  $\rightarrow k$



$$\text{Rate} = k [\text{H}_2\text{O}_2]$$

Order  $\rightarrow 1$

Rate constant  $\rightarrow k$

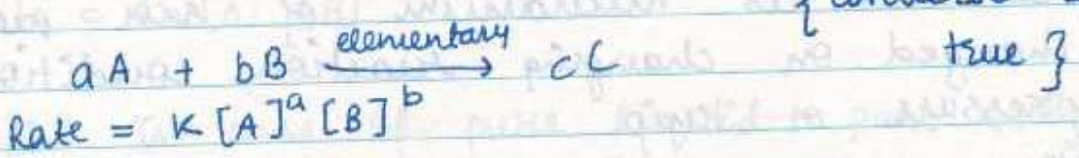


$$\text{Rate} = k_1 \frac{[\text{NH}_3]}{1 + k_2 [\text{NH}_3]}$$

Order  $\rightarrow -\text{N.D.}$

Rate constant  $\rightarrow \text{N.D.}$

- It is not necessary that rate constant as well as order of rxn is defined for every rxn.
- In case of elementary rxn, coeff. of reactants becomes equal to its order



### \* Physical significance of 'K'

- larger the value of 'K' faster will be the rxn. Under given catalytic conditions, 'K' can be changed only by changing temperature.

#### • Units of 'K' :-



$$\text{Rate} = K [A]^n$$

$$K = M^{1-n} s^{-1}$$

Units of 'K' will depend on order of rxn.

if  $n=0 \Rightarrow$  zero order  $\Rightarrow K \rightarrow M s^{-1}$

if  $n=1 \Rightarrow$  first order  $\Rightarrow K \rightarrow s^{-1}$

if  $n=2 \Rightarrow$  second order  $\Rightarrow K \rightarrow M^{-1} s^{-1}$

#### • Various types of 'K' :-



$$ROR = -\frac{1}{a} \frac{d[A]}{dt} = -\frac{1}{b} \frac{d[B]}{dt} = \frac{1}{c} \frac{d[C]}{dt} = \frac{1}{d} \frac{d[D]}{dt} = K_8 [A]^\alpha [B]^\beta$$

$k_r \rightarrow$  rate constant of Rxn

$$-\frac{d[A]}{dt} = a k_r [A]^\alpha [B]^\beta = k_A [A]^\alpha [B]^\beta$$

$$-\frac{d[B]}{dt} = b k_r [A]^\alpha [B]^\beta = k_B [A]^\alpha [B]^\beta$$

where,

$k_A, k_B \rightarrow$  rate of disappearance consts. w.r.t A & B

$$\frac{d[c]}{dt} = c k_r [A]^\alpha [B]^\beta = k_c [A]^\alpha [B]^\beta$$

$$\frac{d[D]}{dt} = d k_r [A]^\alpha [B]^\beta = k_D [A]^\alpha [B]^\beta$$

where,

$k_c, k_D \rightarrow$  rate of app. const.

$$\Rightarrow k_r = \frac{k_A}{a} = \frac{k_B}{b} = \frac{k_c}{c} = \frac{k_D}{d}$$

Note:

- If nothing is mentioned than we will take it as

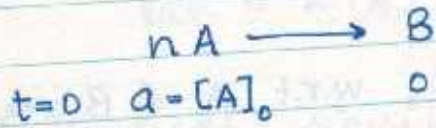


- Units of K helps in determination of order of reaction in numerical problems.

## \* Analysis of Various Orders of Reactions $\Rightarrow$

(Using Integrated Rate Law Expressions)

### i. Zero Order $\Rightarrow$



$$t=t \quad a-x = [A]_0 - \frac{x}{n}$$

$$RDR = -\frac{d[A]}{dt} = \frac{d[B]}{dt} = k_r$$

$$\Rightarrow -\frac{d[A]}{dt} = n k_r = K_A$$

$$\Rightarrow \int_{[A]_0}^{[A]_t} d[A] = \int_0^t K_A dt$$

$$\Rightarrow [A]_t - [A]_0 = K_A t$$

$$\Rightarrow x = K_A t$$

$$\Rightarrow [A]_t = [A]_0 - K_A t$$

$$P_{A_t} = P_{A_0} - K'_A t$$

R

- Half life ( $t_{1/2}$ ) :  $\rightarrow$  It is the time taken to react half of reactant.

$$[A]_t = \frac{[A]_0}{2}$$

$$[A]_t = [A]_0 - K_A t_{1/2}$$

$$\Rightarrow t_{1/2} = \frac{[A]_0}{2K_A} \quad t_{1/2} \propto [A]_0$$

- Completion time ( $t_{\text{comp}}$ )

$$[A]_t = 0$$

$$\Rightarrow t_{\text{comp}} = \frac{[A]_0}{K_A}$$

- Reln b/w  $t_{1/2}$  &  $t_{\text{comp}}$  :

$$t_{\text{comp}} = 2 t_{1/2}$$

- Characteristics of zero order Rxn :

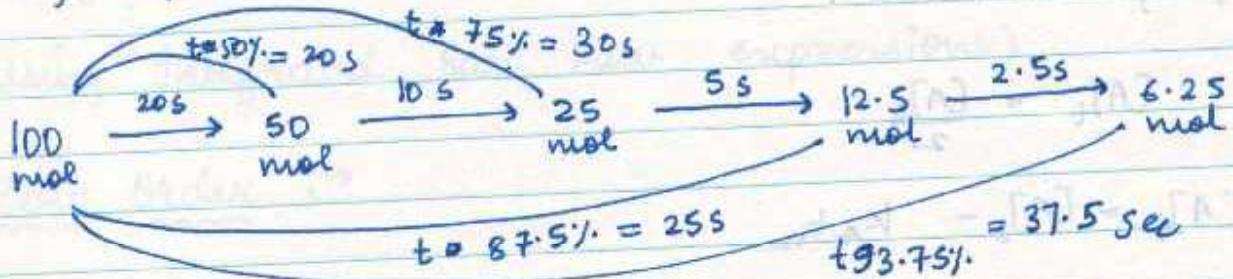
i] Units of  $K \rightarrow \text{Ms}^{-1}$

ii] Conc. of reactant decreases linearly with time

$$[A]_t = [A]_0 - K_A t$$

$$y = c + mx$$

iii) Half life depends on initial conc. of reactant



iv)  $t_{\text{comp}} = \frac{[\text{A}]_0}{k_A}$  { it gets completed in finite time }

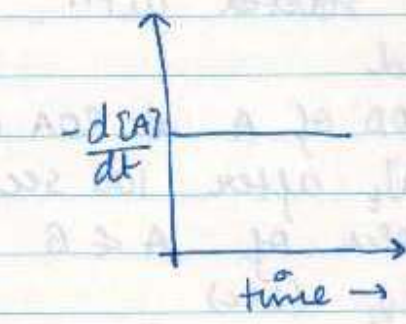
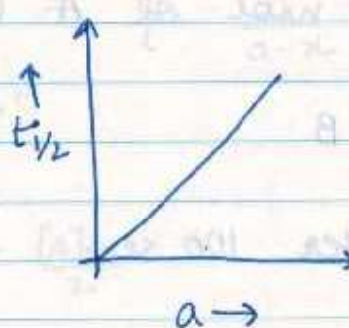
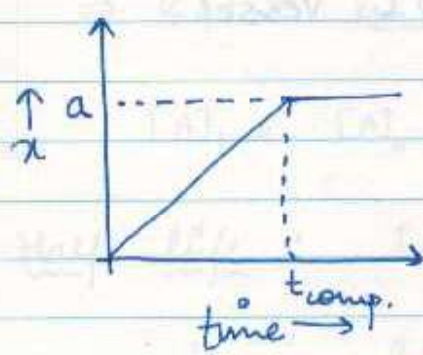
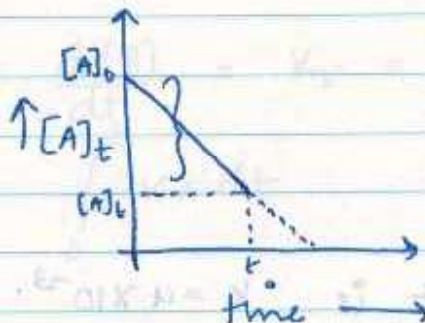
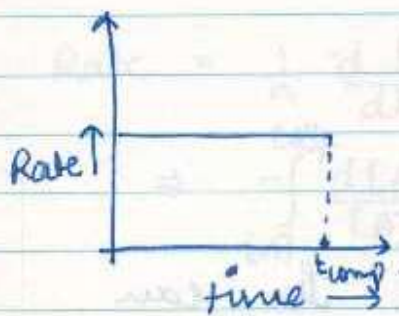
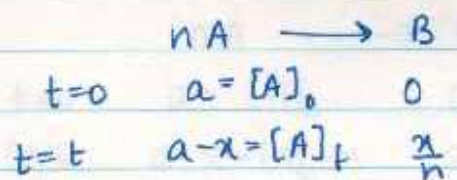
$t_{\text{comp}}$  is also proportional to initial conc.

v) Conc. of reactant in zero order rxn, after equal interval of time constitute an AP with C.D. ( $-k_A t$ ).

time (t)	$[A]_t$	$0$	$t$	$2t$	$3t$	Rate Constant
		$[A]_0$	$[A]_0 - k_A t$	$[A]_0 - 2k_A t$	$[A]_0 - 3k_A t$	
			$-k_A t$	$-k_A t$	$-k_A t$	

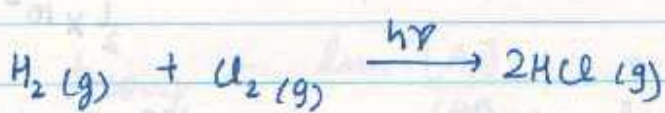
\* vi) Zero order rxn must be complex

• Graphical form :-



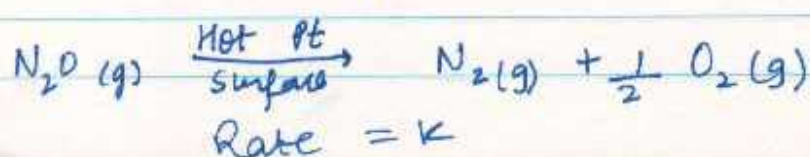
• Examples of zero order :-

i) Photochemical Rxn



$$\text{Rate} = k$$

ii) Surface catalysed Rxn





## 2. First Order Reaction



$$t=0 \quad a = [A]_0 \quad 0$$

$$t=t \quad a-x = [A]_0 - x \quad x$$

$$\text{Rate} = \frac{1}{n} \frac{d[A]}{dt} = \frac{d[B]}{dt} = k_r = k_A [A]_t$$

$$\Rightarrow - \int_{[A]_0}^{[A]_t} \frac{d[A]}{[A]} = \int_0^t k_A dt$$

$$\Rightarrow k_A t = \ln \frac{[A]_0}{[A]_t} = \ln \frac{a}{a-x}$$

$$[A]_t = [A]_0 e^{-k_A t}$$

$$\text{Half life} : [A]_t = \frac{[A]_0}{2}$$

$$t_{1/2} = \frac{\ln 2}{k_A}$$

• Half life will be independent of initial conc.

completion time :

$$t_{\text{comp}} = \lim_{[A]_t = 0} \frac{\ln [A]_0}{[A]_t} = \infty$$

• 1<sup>st</sup> order Rxn will never get completed in finite time

for numerical purpose :

a) If % of reactant reacted is given

$$K_A t = \ln \frac{[A]_0}{[A]_t} = \ln \frac{a}{a-x}$$

$$\left. \begin{array}{l} K_A t_{20\%} = \ln \frac{100}{80} \\ K_A t_{40\%} = \ln \frac{100}{60} \end{array} \right\} K_A t_{x\%} = \ln \frac{100}{100-x}$$

b) If fraction of reactant reacted is given

$$\left. \begin{array}{l} K t_{1/3} = \ln \frac{1}{1-1/3} \\ K t_{1/7} = \ln \frac{1}{1-1/7} \end{array} \right\} K t_{1/n} = \ln \frac{1}{1-1/n}$$

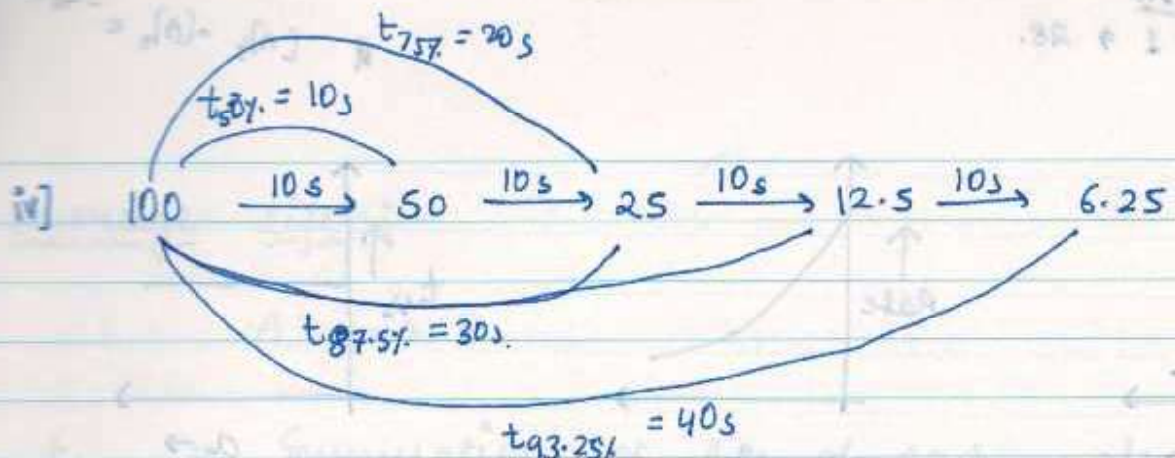
• Characteristics of 1<sup>st</sup> Order Rxn :

i) Units of K  $\rightarrow$  s<sup>-1</sup>

ii) conc. of reactant  $\downarrow$  exponentially with time

$$[A]_t = [A]_0 e^{-K_A t}$$

$$iii) t_{1/2} = \frac{\ln 2}{K_A}$$



v)  $t_{\text{completion}} = \infty$ .

1<sup>st</sup> order reactions never get completed in finite time.

vi) Concentration of reactants after equal intervals of time constitute a G.P. with common ratio  $e^{-k_A t}$ . It means in equal intervals of time, equal % of reactant reacts.

time (t)	0	$t$	$2t$
$[A]_t$	$[A]_0$	$[A]_0 e^{-k_A t}$	$[A]_0 e^{-2k_A t}$
		$e^{-k_A t}$	$e^{-2k_A t}$

vii) Average life ( $t_{\text{avg}}$ )  $= \frac{1}{k_A} = \frac{t_{1/2}}{\ln 2} = 1.44 t_{1/2}$

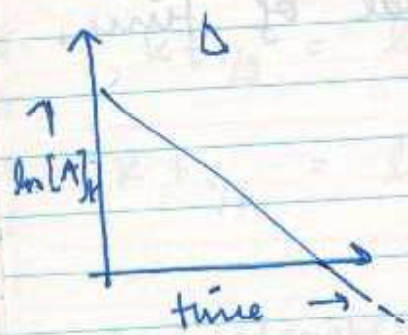
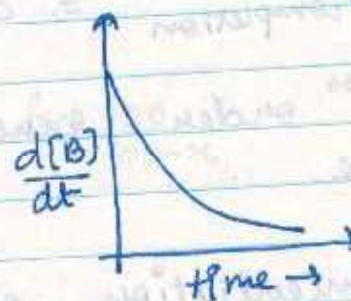
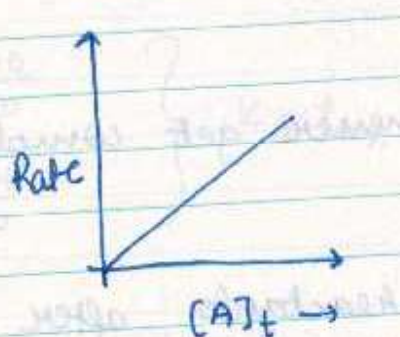
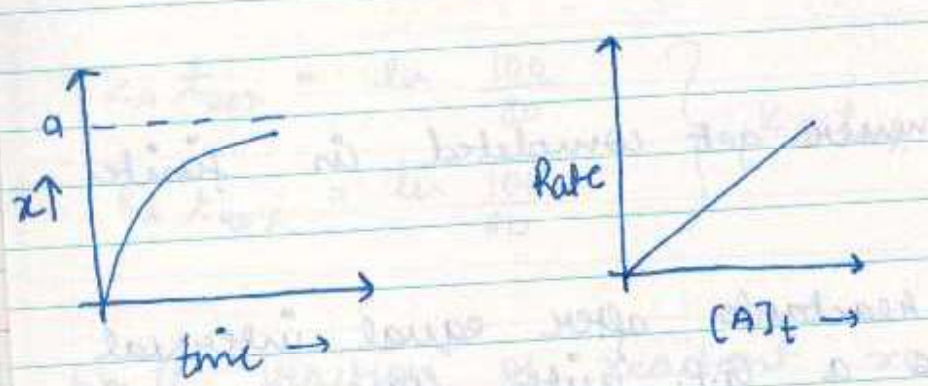
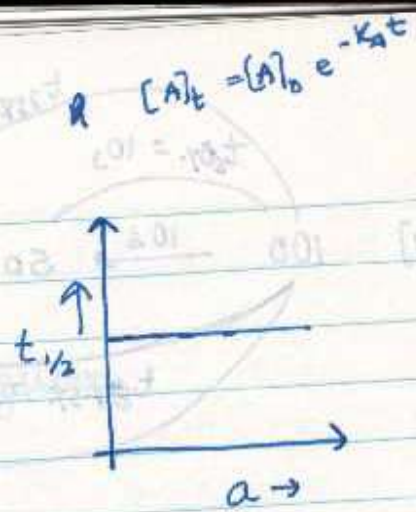
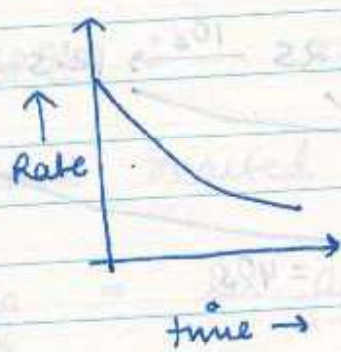
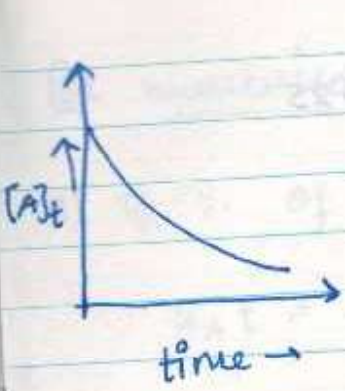
- Graphical Analysis



$$t=0 \quad a=[A]_0$$

$$t=t \quad a-x=[A]_t \quad x$$

$$\frac{H.W.}{Ex 1} \Rightarrow 28.$$



• Average life :



$$\text{t}_{\text{avg.}} = \frac{\text{Summation of life of each molecule}}{\text{Total no. of molecules}}$$

If we take time interval b/w  $t$  to  $t + dt$

$$-\frac{d[A]}{dt} = k_A [A]_t$$

$$-d[A] = k_A [A]_t dt$$

$$\text{t}_{\text{avg.}} = \int_0^{\infty} \frac{k_A [A]_t dt}{[A]_0} = \int_0^{\infty} \frac{k_A [A]_0 t e^{-k_A t} dt}{[A]_0}$$

$$(t_{\text{avg.}})_I = \frac{1}{k_A}$$

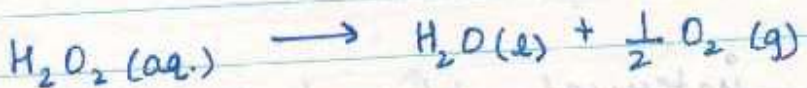
For zero order,

$$-\frac{d[A]}{dt} = k_g = k_A$$

$$\text{t}_{\text{avg.}} = \int_0^t \frac{k_A dt}{[A]_0} = \frac{k_A t^2}{2 [A]_0}$$

Examples of First Order Rxn:

- 1) All radioactive disintegration follows first order kinetics.
- 2) Decomposition of  $H_2O_2$



$$\text{Rate} = k [H_2O_2]$$

Q. For 1 order Rxn P.T.

$$\text{i}) T_{75\%} = 2 t_{50\%}$$

$$\text{ii}) t_{87.5\%} = 3 t_{50\%}$$

$$\text{iii}) t_{99.9\%} = 10 t_{50\%}$$

$$\rightarrow \frac{1}{K_A} \times \ln \frac{100}{25} - ① \quad | \quad \frac{1}{K_A} \ln \frac{100}{50} - ②$$

from ① & ②

$$t_{75} = 2 t_{50}$$

$$t_{87.5\%} = \frac{1}{K_A} \ln 8 = 3 t_{50}$$

$$t_{99.9\%} = \frac{1}{K_A} \ln \frac{100}{0.1} = 10 t_{50}$$

Q. For a 1st order Rxn  $\text{R}_{\text{Rxn}}$  at  $t_1 = 10 \text{ min}$  is  $2 \times 10^{-3} \text{ ms}^{-1}$   
 & at  $t_2 = 20 \text{ min}$  is  $4 \times 10^{-3} \text{ ms}^{-1}$  find  $K$ .

$$\rightarrow \text{R}_{\text{Rxn}} = K_A [A]_t$$

$$= K_A [A]_0 e^{-K_A t}$$

$$\frac{1}{2} = \frac{e^{-K_A 100}}{e^{-K_A 200}}$$

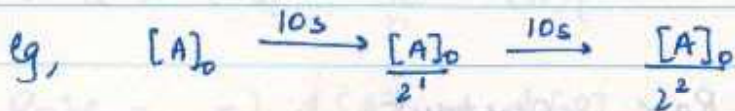
$$\Rightarrow K_A \times 600 = -\frac{\ln 2}{600} = 4 \text{ s}^{-1}$$

$$\frac{2 \ln 2}{600} = \frac{1.4}{600}$$

for numerical purpose

$\frac{[A]_t}{[A]_0}$  = fraction of reactant left

$$1 - \frac{[A]_t}{[A]_0} = \text{" " " reacted.}$$



$$[A]_t = \frac{[A]_0}{2^n}$$

where  $n = \text{no. of half lives}$

$$\frac{[A]_t}{[A]_0} = \frac{1}{2^n}$$

$$\rightarrow 1 - \frac{1}{2^n} = 63/64 \times 100$$

### Pseudo Order Rxn:

If more than one reactants are present & one of them is in large excess then during rxn conc. of reactant which is present in excess qty. almost becomes constant. In such a case its conc. is merged with  $k$  & order w.r.t. other reactant (lesser) is called pseudo order.

	$A + B \longrightarrow C$		
$t=0$	1000M	1M	0
$t=t_1$	999.5M	0.5M	0.5M
$t=t_2$	999.25M	0.25M	0.75M

$$\text{rate} = k_p [A]^1 [B]^1$$

$$\frac{r_1}{r_2} = \frac{k_p \times 1000 \times 1}{k_p \times 999.5 \times 0.5} = \frac{2}{1}$$

$$\therefore \text{rate} = k' [B]$$

where,

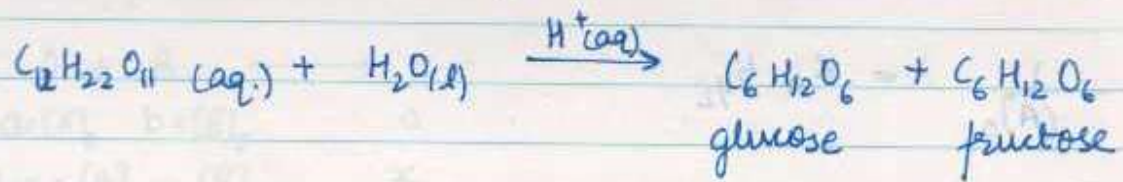
$$k' = k[A] = \text{Pseudo Rate constant}$$

$$k't = \ln \frac{[B]_0}{[B]_t}$$

$$\therefore k' = k[A]$$

(Rate constant for rxn.)

e.g. inversion of cane sugar is pseudo I order rxn



$$\text{rate} = k [H_2O][H^+][C_{12}H_{22}O_{11}]$$

$$\text{rate} = k' [C_{12}H_{22}O_{11}]$$

$$k' = k [H_2O][H^+]$$

↓  
Pseudo rate constant

$$k't = \ln \frac{[C_{12}H_{22}O_{11}]_0}{[C_{12}H_{22}O_{11}]_t}$$

### 3. Second Order Rxn:

case I: only one reactant is present



$$t=0 \quad a = [A]_0 \quad 0$$

$$t=t \quad a-n = [A]_t \quad \frac{x}{n} = [B]_t$$

$$\text{Rate} = -\frac{1}{n} \frac{d[A]}{dt} = \frac{d[B]}{dt} = k_2 [A]_t^2$$

$$\left. \frac{-d[A]}{[A]^2} \right|_{(A)_0} = k_A \int_0^t dt$$

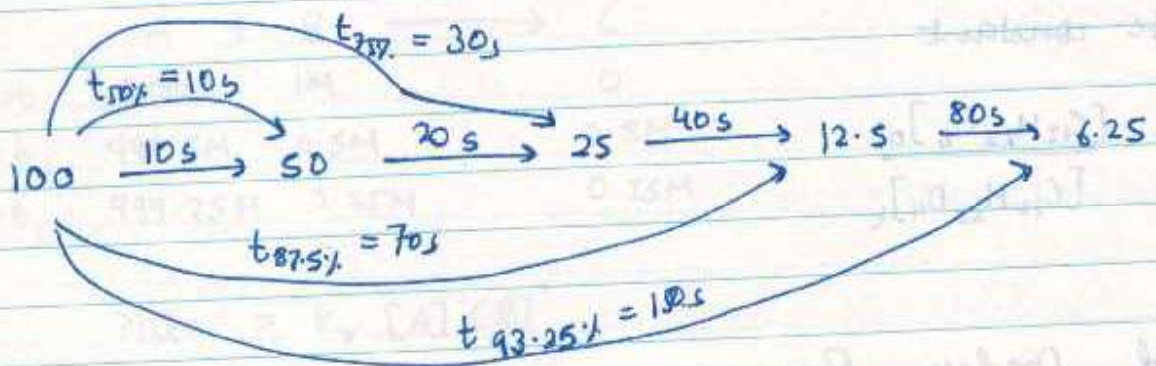
$$\Rightarrow -\left[ \frac{1}{[A]_t} - \frac{1}{[A]_0} \right] = k_A t$$

Half life :

$$\frac{1}{[A]_0} - \frac{1}{[A]_t} = k_A t_{1/2}$$

$$\Rightarrow t_{1/2} = \frac{1}{k_A [A]_0}$$

$$\Rightarrow t_{1/2} \propto \frac{1}{[A]_0}$$

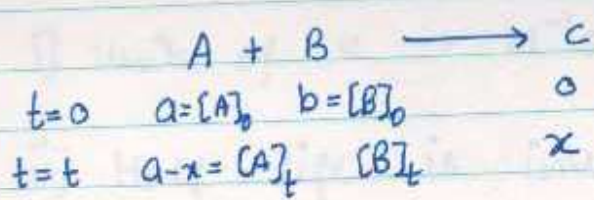


Completion time :

$$[A]_f = 0$$

$t_{comp} = \infty$

Case II : If more than one reactants are present



$$\text{Rate} = -\frac{d[A]}{dt} = -\frac{d[B]}{dt} = \frac{d[C]}{dt} = k_r [A]_t [B]_t$$

$$-\frac{d[A]}{dt} = k_r [A]_t [B]_t$$

$$\Rightarrow -\frac{d(a-x)}{dt} = k_r (a-x) (b-x)$$

$$\Rightarrow \int_{0}^{x} \frac{dx}{(a-x)(b-x)} = \int_{0}^t k_r dt$$

$$\Rightarrow \frac{1}{a-b} \int_{0}^{x} \frac{(a-x)-(b-x)}{(a-x)(b-x)} = \int_{0}^t k_r dt$$

$$\Rightarrow \frac{1}{a-b} \left\{ \int_{0}^{x} \frac{dx}{(b-x)} - \int_{0}^{x} \frac{dx}{(a-x)} \right\} = k_r t$$

$$\Rightarrow \frac{1}{a-b} \left\{ -\ln(b-x) + \ln(a-x) \right\}_{0}^{x} = k_r t$$

$$\Rightarrow \frac{1}{a-b} \left\{ \ln \frac{a-x}{b-x} - \ln \frac{a}{b} \right\} = k_r t$$

$$\Rightarrow \frac{1}{a-b} \left[ \ln \frac{(a-x)b}{(b-x)a} \right] = k_r t$$











If  $k_{A_1} \approx k_{A_2}$   
 $\Rightarrow$  Rate is of zero order

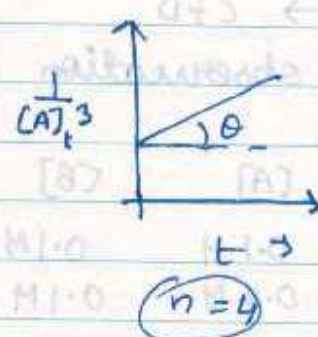
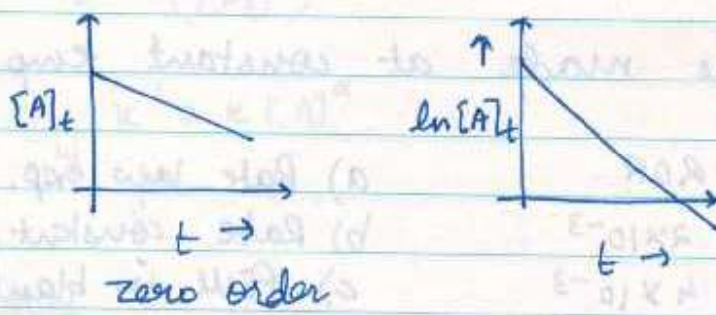
If  $k_{A_1} \neq k_{A_2}$   
we assume order = 1

$$k_{A_1} t_1 = \ln \frac{[A]_0}{[A]_{t_1}}$$

$$k_{A_2} t_2 = \ln \frac{[A]_0}{[A]_{t_2}}$$

If  $k_{A_1} \approx k_{A_2}$   
 $\Rightarrow$  Rate is of first order.

## 2. Graphical Method :



## 3. Half-life Method :

$$t_{1/2} \propto a^{1-n} \propto p^{1-n}$$

$$\frac{(t_{1/2})_I}{(t_{1/2})_{II}} = \left(\frac{a_I}{a_{II}}\right)^{1-n} = \left(\frac{P_{A_I}}{P_{A_{II}}}\right)^{1-n}$$



## 5. Isolation Method :

In this method, when more than one reactants are present then one of the reactant is taken in large excess as a result its conc. remains almost constant during ran & order of ran w.r.t. other reactant is calculated.



$$\text{Rate} = k [A]^{\alpha} [B]^{\beta}$$

$$\text{if } [A] \gg [B]$$

$$\text{Rate} = k' [B]^{\beta}$$

$$\frac{r_1}{r_2} = \left( \frac{[B_1]}{[B_2]} \right)^{\beta}$$

$$k' = k [A]^{\alpha}$$

" for  $\alpha'$ .

H.W

DPP # 9, 10

Ex. 1: 29-36

R.B.: 1, 2, 8, 13-15, 16, 20,

Ex. 2: 48, 49, 52, 53



Q. find  $K$  for,



(t)	0	t	$\infty$
(P)	$P_0$	$P_t$	$P_\infty$

$$P_0 + (n-1)x = P_t \quad \frac{P_t - P_0}{(n-1)}$$

$$\rightarrow K_A t = \ln \frac{P_0}{P_0 - \left[ \frac{P_t - P_0}{(n-1)} \right]}$$

$$= \ln \frac{P_\infty - P_0}{P_\infty - P_t}$$

B. Progress of decomposition of  $\text{NH}_4\text{NO}_2$  (aq) is obs. by total pressure



time (s)	0	20	$\infty$	find	$t_{1/2}$
P <sub>total</sub>	50	80	90		

$$\rightarrow K_A = \frac{1}{20} \times \ln \frac{80}{50} \approx 4$$

$$\underline{M-1} \quad = \frac{\ln 2}{10}$$

$$t_{1/2} = 10 \text{ s}$$

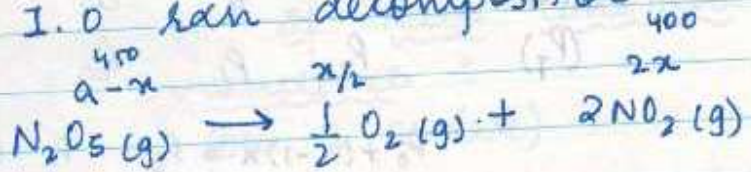
~~M-II~~

$$\alpha \propto 30 \quad K_A t = \ln \frac{a}{a-x} = \ln \frac{40}{10}$$

$$a \propto 40$$

$$a = 40 \quad x = a(1 - e^{-K_A t})$$

For I.O can decomposition



Initially the total pressure is found to be 650 torr & after a very long time total pressure is found to be 1550 torr. If after 4 min from start of Rxn  $P_{O_2} = 100$  torr calculate  $t_{1/2}$  [  $\ln 3 = 1.1$ ,  $\ln 2 = 0.7$  ]

$$\begin{aligned}\rightarrow K_A t = \ln \frac{900}{200} &= 2\ln 3 - \ln 2 & K = \frac{1}{4} \times \ln \frac{650}{400} \\ &= 2 \cdot 2 - 0.7 = \frac{1.5}{4} & \text{13}\end{aligned}$$

7  $t_{1/2} = \frac{0.7 \times 4}{1.5} = \frac{28}{15} = \boxed{2}$

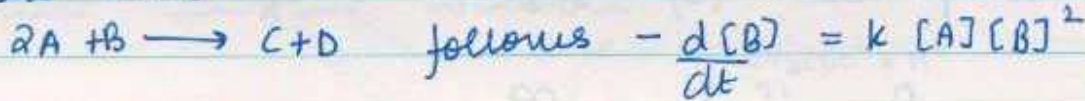
If  $P_t$  is given at initial & after long time then we can check it for presence of inert gas.

Q. For a Rxn  $A(g) \xrightarrow{n} B(g)$  the rate constant is  $6.93 \times 10^{-4} \text{ s}^{-1}$ . The Rxn is performed at constant pressure & temp. of  $24.63^\circ\text{C}$  & 300 K. Starting with 1 mol of pure A. If conc. of B after 2000 sec is  $\frac{3}{3.25} \text{ M}$ . Then calculate 'n'.

$$\begin{aligned}\rightarrow \frac{nA}{V} &= \frac{3}{3.25} & 4 = \frac{a}{a-x} & 6.93 \times 10^{-4} \times 2 \times 10^3 = \ln \\ & n - nx = 1 & x = \frac{3}{4} & 1.386 = \ln 4\end{aligned}$$

14

Q. The rxn



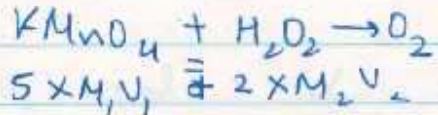
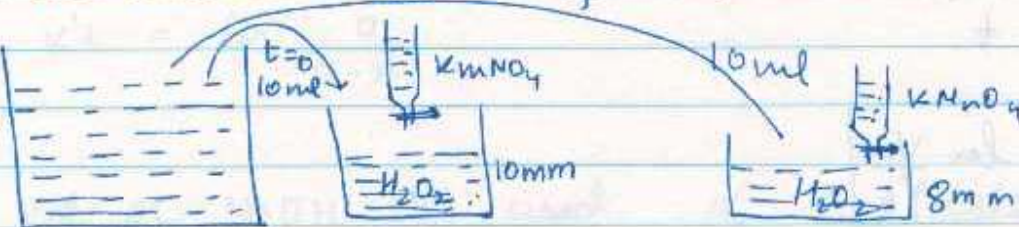
Find (x+y)

S. No.	$[A]_0 \times 10^{-4}$ M	$(B)_{0.2} \times 10^{-4}$ M	$t_{1/2}$ (sec)	$\frac{t_{1/2}}{[B]} \text{ sec}$
1	500	2	10	
2	500	4	x	= $10 \times 1/5$
3	2	250	8	
4	4	500	y	= $8 \times 1/2$

$$\left(\frac{1}{2}\right)^x$$

ii] In terms of decomposition of  $H_2O_2$  (aq.) :

In this method a sol<sup>n</sup> of  $H_2O_2$  is prepared and equal amount of sample or aliquot is taken from  $H_2O_2$  sol<sup>n</sup> & after certain interval of time, it is titrated against standard sol<sup>n</sup> of  $KMnO_4$  to determine value of 'k'.

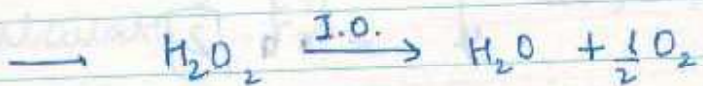


g. For decomposition of  $H_2O_2$

Time (t) 0 t  $\infty$

Vol. of KMnO<sub>4</sub> used (in ml)  $V_0$   $V_t$   $V_\infty$

Find  $t_{1/2}$ .



$t=0 \quad a=1 \quad 0 \quad 0$   
 $t=t \quad a-x \quad x \quad \frac{x}{2}$

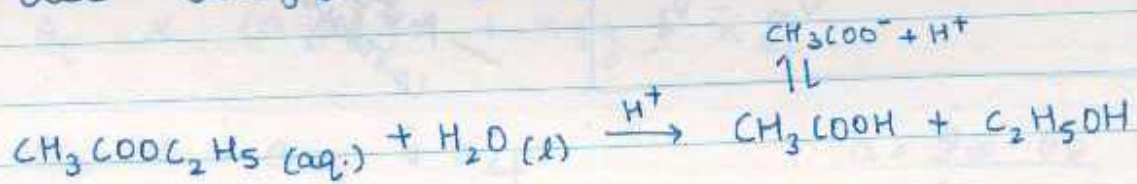
$$K_A t = \ln \frac{a}{a-x}$$

• Vol. of KMnO<sub>4</sub> ( $V_0$ ) used  $\propto a$   
initially

• Vol. of KMnO<sub>4</sub> ( $V_t$ ) used  $\propto a-x$   
after time t

$$\therefore K_A t = \ln \frac{V_0}{V_t}$$

iii] Acid catalysed hydrolysis of ester :→



$t=0$	$a$	$0$	$0$
$t=t$	$a-x$	$x$	$x$
$t=\infty$	$0$	$a$	$a$

Q. For hydrolysis of ester

time ( $t$ )	0	$t$	$\infty$
Vol. of NaOH used (in ml)	$V_0$	$V_t$	$V_\infty$

$$\rightarrow \text{Rate} = k [H^+] [H_2O] [CH_3COOC_2H_5]$$

$$\Rightarrow \text{Rate} = k' [CH_3COOC_2H_5]$$

$$k't = \ln \frac{a}{a-x}$$

Vol. of NaOH ( $V_b$ ) used  $\propto$  C (catalyst)  
initially

Vol. of NaOH ( $V_t$ ) used  $\propto$  C + x  
after time  $t$

Vol. of NaOH ( $V_\infty$ ) used  $\propto$  C + a  
after long time



$$\theta_0 \propto \alpha \times r_s$$

$$\theta_t \propto (\alpha - x) r_s + x r_g + x r_f$$

$$\theta_{\infty} \propto x r_g + x r_f$$

$$\alpha = \frac{\theta_{\infty} - \theta_0}{r_g - r_f - r_s}$$

$$K_A t = \ln \frac{a}{a-n}$$

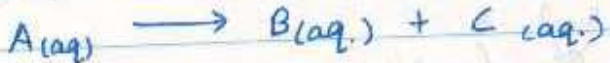
$$= \ln \frac{\theta_{\infty} - \theta_0}{\theta_{\infty} - \theta_t}$$

$$n = \frac{\theta_t - \theta_0}{r_g - r_f - r_s}$$

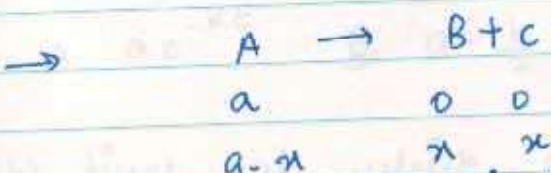
Note :

- Resemic mixture is optically inactive, it means angle of rotation will be zero.

Q. For a rxn



Starting with only A(aq), optical rotation after 10 min & 20 min is obs. to be  $33^\circ$  &  $18^\circ$  when initial optical rotation was  $48^\circ$ . If specific optical rotation for A, B & C are  $20^\circ$ ,  $30^\circ$  &  $-40^\circ$  calculate the time (in min) when a resemic mixture will be formed.



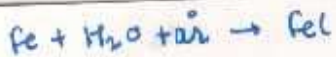
$$K_A 10 = \ln \frac{a}{a-n}$$

$$K_A 20 = \ln \frac{a}{a-n-y}$$

$$\begin{aligned} (a-n) \times 20^\circ + 0 \times 30^\circ - n \times 40^\circ &= 33^\circ \\ (a-n-y) \times 20^\circ + y \times 30^\circ + x \times 30^\circ - (x+y) \times 40^\circ &= 18^\circ \end{aligned}$$

*32 min*

$$\frac{1}{2} = \frac{\ln (a-(n+y))}{\ln (a-n)}$$



$$x_A t = \ln \frac{a}{a-x}$$

200

### v) Thickness of Oxide layer formation :

If oxide formation follows 1<sup>st</sup> order kinetics

time       $t$        $\infty$       Find  $k$ .

Thickness of layer	$Z_t$	$Z_\infty$	$k t = \ln \frac{Z_\infty}{Z_\infty - Z_t}$
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### vi) Bacteria growth :

If bacteria growth follows 1<sup>st</sup> order kinetics



$t=0$        $b$

$t=t$        $b+x$

$$\begin{aligned} \frac{dB}{dt} &= k B \Rightarrow \int_b^{b+x} \frac{dB}{B} = \int_0^t k dt \\ &= \ln \frac{B}{b} \Big|_b^{b+x} = k t \\ &= \ln \frac{b+x}{b} = k t \end{aligned}$$

Q. After how much time no. of bacteria will become double of its initial value.

$$\rightarrow k t = \ln 2$$

$$\Rightarrow t = \frac{\ln 2}{k}$$

Q. For a Rxn



Kinetics is determined by analysis of partial pressure of gases at different instants. Calculate  $a+b+c$ .

$$P \text{ (mm of Hg)} \quad t=0 \quad t=30 \text{ min} \quad t=60 \text{ min}$$

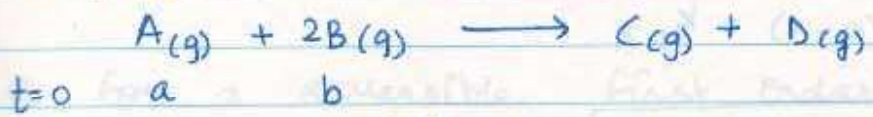
$P_A$	20	10	$a = 5$
$P_B$	10000	NA	NA
$P_C$	0	NA	15
$P_D$	0	b	NA

Order of Rxn with respect to A = C = 1,  $a = 5, b = 2$

$$\rightarrow K_p = K' P^A \quad (8) \quad K \times t = \ln \frac{20}{10} \Rightarrow K = \frac{\ln 2}{30}$$

$$\frac{\ln 2}{30} \times \frac{60}{2} = \ln \frac{20}{a}$$

Q. Consider the following rxn occurring at constant volume



$$\text{Rate} = K[B]$$

a) conc. of 'A' at time t is given as

- A.  $ae^{-kt}$     B.  $a - \frac{b}{2}(1-e^{-2kt})$     C.  $a - kt$     D.  $a - b(1-e^{-kt})$

b) time at which conc. of A reduces to half

- A.  $\frac{a}{2k}$     B.  $\frac{a}{k}$     C.  $\frac{1}{2k} \ln \frac{b}{a}$     D.  $\frac{\ln 2}{k}$

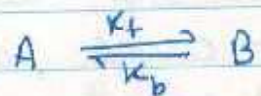
c) For what cond.  $(t_{1/2})_A = (t_{1/2})_B$

- A.  $\frac{a}{b} = \frac{1}{2}$     B.  $\frac{a}{b} = 1$     C. None

Note: If more than one reactants are present then half life of l.s. will be half life of each.

\* Reversible Elementary Rxn:

i) First order opposed by first order:



$$K_{eq} = \frac{k_f}{Rb} = \frac{[B]_{eq}}{[A]_{eq}} = \frac{x_e}{a-x_e}$$

$t=0$	$a$	$0$
$t=t$	$a-x$	$x$
$t=t_{eq}$	$a-x_e$	$x_e$

$$-\frac{d[A]}{dt} = k_f [A] - k_b [B]$$

$$\Rightarrow -\frac{d(a-x)}{dt} = k_f (a-x) - k_b x$$

$$\frac{dx}{dt} = a k_f - x (k_f + k_b)$$

At equilibrium,

$$-\frac{d[A]}{dt} = 0 = k_f (a-x_e) - k_b x_e$$

$$a k_f = x_e (k_f + k_b)$$

$$\Rightarrow \frac{dx}{dt} = (k_f + k_b)(x_e - x)$$

$$\Rightarrow \int_0^x \frac{dx}{x_e - x} = \int_0^t (k_f + k_b) dt$$

$$\Rightarrow -[\ln(x_e - x)]_0^\infty = (k_f + k_b)t$$

$$\Rightarrow (k_f + k_b)t = \ln \frac{x_e}{x_e - x}$$

where,

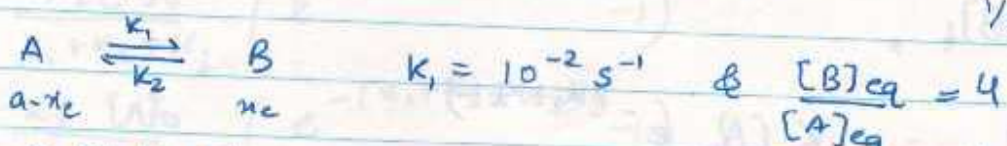
$x_e \rightarrow$  conc. of B at eq.

Q. calculate time at which conc. of B becomes half of its value at equilibrium.

$$\rightarrow (k_f + k_b)t = \ln 2$$

$$\Rightarrow t = \frac{\ln 2}{(k_f + k_b)}$$

Q. for a reversible first order reaction



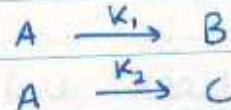
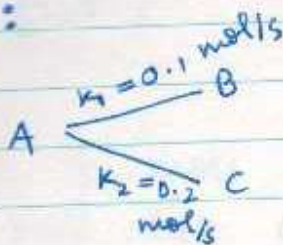
If  $[A]_0 = 0.01 \text{ M L}^{-1}$  &  $[B]_0 = 0$   
what will be the conc. of B after 30 sec.

$$\rightarrow 0.01 \times 10^{-2} = [B]_{eq} (k_f + k_b)$$

$$\left| \begin{array}{l} \frac{10^{-4}}{[B]_{eq}} \times \frac{100 \times 10^{-4}}{30} = \ln \frac{4/100}{[B]_{eq} - x} \\ \frac{0.15}{4} \end{array} \right. \quad \begin{array}{l} \frac{4/100}{5} \\ \frac{4}{4-100x} \end{array}$$

1st order parallel or side rxn:

I :



hyp.

$$-\frac{d[A]}{dt}_{\text{sum}} = \frac{d[B]}{dt} = k_1 [A]$$

$$-\frac{d[A]}{dt}_{\text{sum}} = \frac{d[C]}{dt} = k_2 [A]$$

$$-\frac{d[A]}{dt} = \left( -\frac{d[A]}{dt} \right)_I + \left( -\frac{d[A]}{dt} \right)_{II}$$

$$-\frac{d[A]}{dt} = (k_1 + k_2) [A]$$

$$\Rightarrow - \int \frac{d[A]}{[A]} = \int (k_1 + k_2) dt$$

$$\Rightarrow [A]_t = [A]_0 e^{-(k_1 + k_2)t}$$

For  $[B]_t$ ,

$$\frac{d[B]}{dt} = k_1 [A]_0 e^{-(k_1 + k_2)t}$$

$$\Rightarrow [B]_t = -\frac{k_1 [A]_0}{k_1 + k_2} \left[ e^{-(k_1 + k_2)t} - 1 \right]$$

Similarly

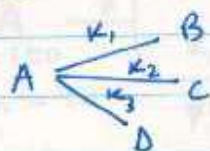
$$[C]_t = \frac{k_2[A]_0}{k_1+k_2} \left( 1 - e^{-(k_1+k_2)t} \right)$$

$$\frac{[B]_t}{[C]_t} = \frac{k_1}{k_2}$$

$$\% \text{ B in product} = \frac{k_1}{k_1+k_2} \times 100$$

$$\% \text{ C in product} = \frac{k_2}{k_1+k_2} \times 100$$

• Case II :



$$-\frac{d[A]}{dt} = (k_1 + k_2 + k_3) [A]$$

$$\Rightarrow [A]_t = [A]_0 e^{-(k_1+k_2+k_3)t}$$

also,

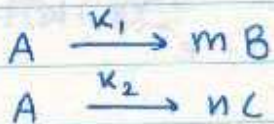
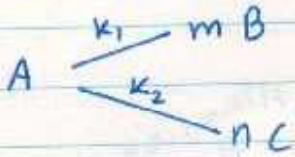
$$[B]_t = -\frac{k_1[A]_0}{k_1+k_2+k_3} \left( e^{-(k_1+k_2+k_3)t} - 1 \right)$$

$$[C]_t = -\frac{k_2[A]_0}{k_1+k_2+k_3} \left( e^{-(k_1+k_2+k_3)t} - 1 \right)$$

$$[D]_t = -\frac{k_3[A]_0}{k_1+k_2+k_3} \left( e^{-(k_1+k_2+k_3)t} - 1 \right)$$

$$\therefore [B]_t : [C]_t : [D]_t = k_1 : k_2 : k_3$$

• Case III :



$$-\frac{d[A]}{dt} \underset{\textcircled{I}}{=} \frac{1}{m} \frac{d[B]}{dt} = k_1 [A]$$

$$-\frac{d[A]}{dt} \underset{\textcircled{II}}{=} \frac{1}{n} \frac{d[C]}{dt} = k_2 [A]$$

$$\Rightarrow -\frac{d[A]}{dt} = (k_1 + k_2) [A]$$

$$\Rightarrow [A]_t = [A]_0 e^{-(k_1 + k_2)t}$$

for  $[B]_t$ ,

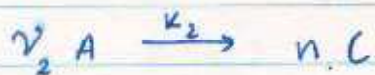
$$\frac{1}{m} \frac{d[B]}{dt} = k_1 [A]_0 e^{-(k_1 + k_2)t}$$

$$\Rightarrow [B]_t = \frac{m k_1 [A]_0}{(k_1 + k_2)} \left( 1 - e^{-(k_1 + k_2)t} \right)$$

$$[C]_t = \frac{n k_2 [A]_0}{(k_1 + k_2)} \left( 1 - e^{-(k_1 + k_2)t} \right)$$

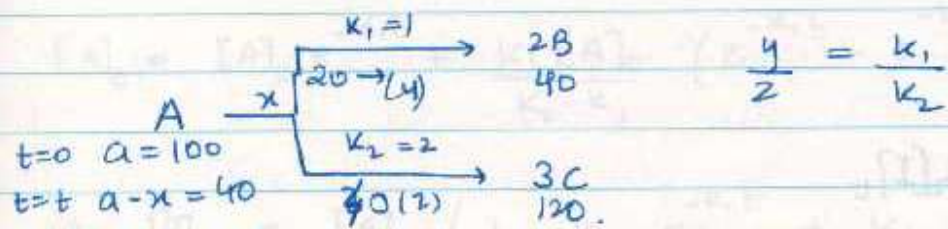
$$\frac{[B]_t}{[C]_t} = \frac{m k_1}{n k_2}$$

• Case IV :

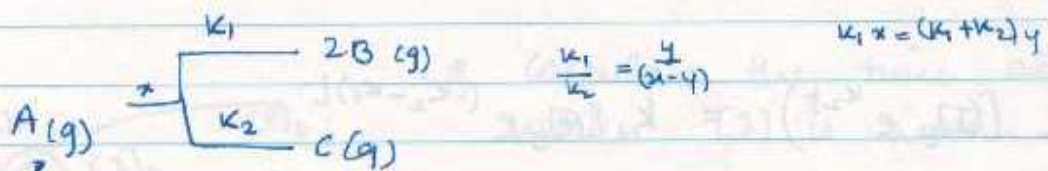


$$-\frac{d[A]}{dt} = (\nu_1 k_1 + \nu_2 k_2) [A]$$

For numerical Purpose,



Q. For following 1 order gaseous rxn.



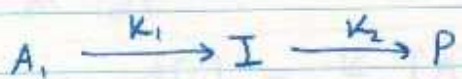
The initial pressure in a container of cap.  $V L$  1 atm. Pressure at  $t = 10 s$  is 1.4 atm & after infinite time it becomes 1.5 atm. Find  $k_1$  &  $k_2$

$$\rightarrow 1-x + \frac{2k_1 x}{k_1+k_2} + \frac{k_2 x}{k_1+k_2} = 1.4$$

$$0 + \frac{2k_1}{k_1+k_2} + \frac{k_2}{k_1+k_2} = 1.5$$

$$\frac{k_1}{k_1+k_2} = 0.5$$

\* consecutive or sequential Rxn:



$$t=0 \quad a \quad 0 \quad 0$$

$$t=t \quad a-x \quad x-y \quad y$$

$$\cdot [A]_t = [A]_0 e^{-k_1 t}$$

$$\cdot \frac{d[I]}{dt} = k_1 [A]_t - k_2 [I]_t$$

$$e^{k_2 t} \frac{d[I]}{dt} + e^{k_2 t} k_2 [I]_t = k_1 [A]_0 e^{(k_2 - k_1) t}$$

$$\Rightarrow \frac{d}{dt} ([I]_t e^{k_2 t}) = k_1 [A]_0 e^{(k_2 - k_1) t}$$

$$\int d([I]_t e^{k_2 t}) = k_1 [A]_0 \int e^{(k_2 - k_1) t} dt$$

$$\Rightarrow [I]_t e^{k_2 t} = \frac{k_1 [A]_0}{(k_2 - k_1)} e^{(k_2 - k_1) t} + C$$

$$\text{At } t=0 \quad [I]_t = 0$$

$$0 = \left( \frac{k_1 [A]_0}{k_2 - k_1} \right) + C$$

$$\Rightarrow [I]_t e^{k_2 t} = \frac{k_1 [A]_0}{k_2 - k_1} (e^{(k_2 - k_1)t} - 1)$$

$$[I]_t = \frac{k_1 [A]_0}{k_2 - k_1} (e^{-k_1 t} - e^{-k_2 t})$$

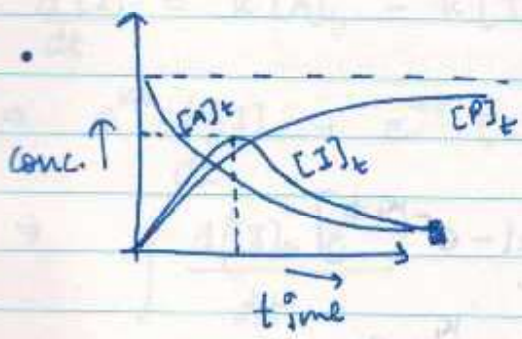
for  $[P]_t$ ,

$$a = (a - x) + (x - y) + y$$

$$[A]_0 = [A]_t + [I]_t + [P]_t$$

$$[A]_0 = [A]_0 e^{-k_1 t} + \frac{k_1 [A]_0}{k_2 - k_1} (e^{-k_1 t} - e^{-k_2 t}) + [P]_t$$

$$\Rightarrow [P]_t = [A]_0 \left( 1 - \frac{k_2}{k_2 - k_1} e^{-k_1 t} + \frac{k_1}{k_2 - k_1} e^{-k_2 t} \right)$$

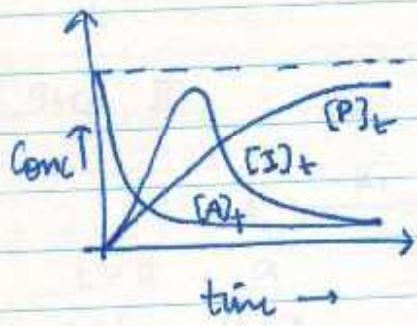
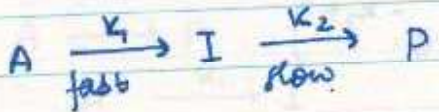


Q. Calculate the time at which  $[I]$  is max.

$$\frac{d[I]_t}{dt} = \frac{k_1 [A]_0}{k_2 - k_1} (-k_1 e^{-k_1 t} + k_2 e^{-k_2 t})$$

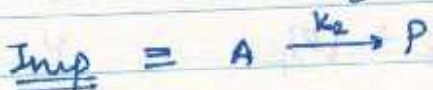
$$\Rightarrow t = \ln \frac{k_2}{k_1} \frac{1}{(k_2 - k_1)}$$

Case I: if  $k_1 \gg k_2$



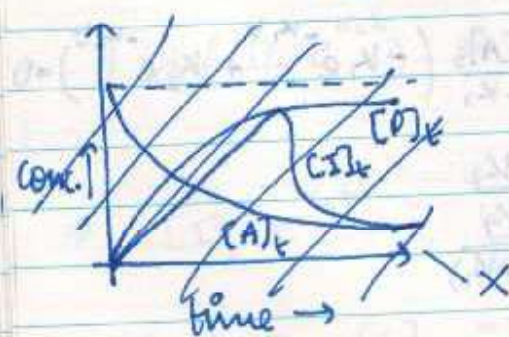
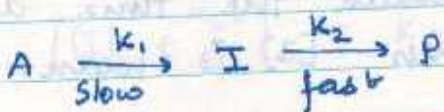
$$[P]_t = [A]_0 \left[ 1 - \frac{k_2}{k_2 - k_1} e^{-k_1 t} + \frac{k_1}{k_2 - k_1} e^{-k_2 t} \right]$$

$$[P]_t = [A]_0 \left( 1 - e^{-k_2 t} \right)$$

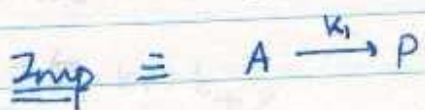


• Conc. of product will depend on slowest step of rxn. From here comes the concept of Rate determining step (RDS). So slowest step will be R.D.S.

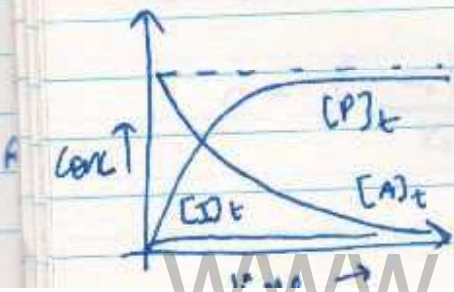
Case II: if  $k_2 \gg k_1$



$$[P]_t = [A]_0 \left[ 1 - e^{-k_1 t} \right]$$



Steady state: It is the state at which rate of formation of intermediate becomes equal to rate of disappearance of intermediate or conc. of I becomes very very small.



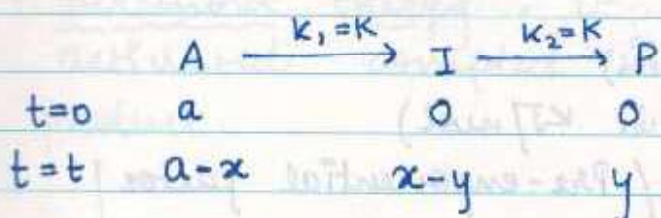
For steady state,  $K_2 \gg K_1$

$$\frac{d[I]}{dt} = 0 = K_1[A]_t - K_2[I]_t$$

$$\Rightarrow [I]_t = \frac{K_1}{K_2} [A]_t$$

at steady state.

Case III: if  $K_1 = K_2 = K$



$$[A]_t = [A]_0 (e^{-kt})$$

$$\frac{d[I]}{dt} = K[A]_t - K[I]_t = K \{ [A]_t - [I]_t \}$$

$$\Rightarrow e^{kt} \frac{d[I]}{dt} + e^{kt} [I]_t K = K[A]_0$$

$$\Rightarrow \int \frac{d([I]_t e^{kt})}{dt} = \int K[A]_0 dt$$

$$\Rightarrow K[A]_0 t = [I]_t e^{kt} + C$$

$$\text{At } t=0, [I]_t = 0$$

$$0 = 0 + C$$

$$[I]_t = K[A]_0 + e^{-kt}$$

## \* Arrhenius Equation :-

It is quantitative relationship proposed by Arrhenius b/w rate constant and temperature.

$$K = A e^{-E_a/RT}$$

where,

$K \rightarrow$  Rate constant / Specific Reaction Rate / Velocity constant

$R \rightarrow$  Gas constant

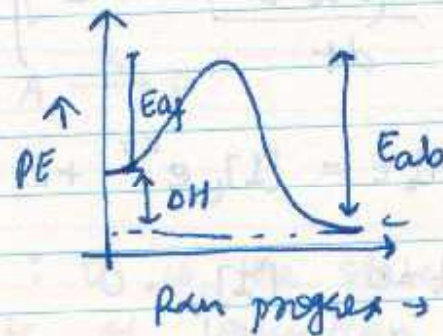
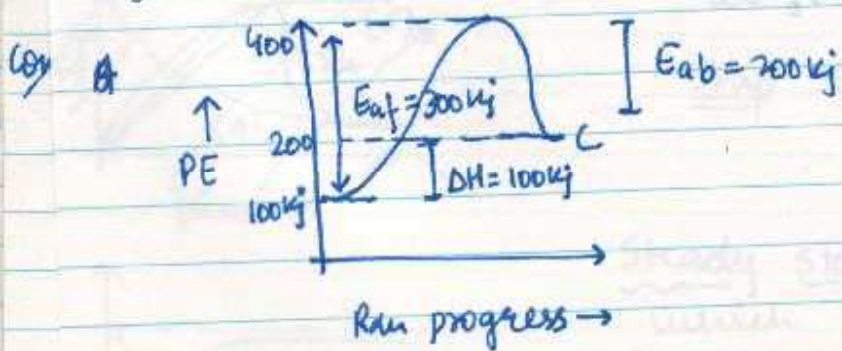
$T \rightarrow$  Temperature (in Kelvin)

$E_a \rightarrow$  Activation Energy (in  $\text{kJ/mol}$ )

$A \rightarrow$  Arrhenius constant / Pre-exponential factor / frequency factor.

(  $A$  &  $E_a$  are considered as temperature independent.

[Although  $A$  &  $E_a$  are considered as depends on temp. but on change in temp. their values change negligibly.]



Core



$$\Delta H = E_{\text{af}} - E_{\text{ai}} = E_p - E_R$$

if  $\Delta H < 0 \Rightarrow$  Rxn is exothermic

$\Delta H > 0 \Rightarrow$  Rxn is endothermic

• Activation Energy: Minimum K.E. possessed by reactant molecules in order to get converted into product.

• Threshold Energy: It is min. P.E. possessed by activated complex in order to get converted to product.

### \* Variation of $K$ :

- For complete elementary rxn (not elementary step)  $E_A > 0$ , but in case of complex rxn  $E_A$  can be -ve or zero.

$$K = A e^{-E_a/RT}$$

$\Rightarrow$  if  $T \uparrow \Rightarrow K \uparrow \Rightarrow$  ROR will increase (in general)

$\Rightarrow$  if  $E_a \uparrow \Rightarrow K \downarrow \Rightarrow$  ROR will decrease

$\Rightarrow$  At  $T=0K \Rightarrow K=0$

$\Rightarrow$  At  $T=\infty \Rightarrow K=A$

Q. On ↑ temp. rate of rxn will always change.

→ False. If  $E_A = 0$ , then  $K = A$  &  
A is independent of temp.

\* Variation of  $K$  with temperature:

$$K = A e^{-E_a/RT}$$

Considering A &  $E_a$  to be temp. independent

$$\ln K = \ln A = -\frac{E_a}{RT}$$

At temp.  $T_1$ ,

$$\ln K_{T_1} = \ln A - \frac{E_a}{RT_1} \quad \text{--- (1)}$$

At temp.  $T_2$ ,

$$\ln K_{T_2} = \ln A - \frac{E_a}{RT_2} \quad \text{--- (2)}$$

$$(1) - (2)$$

$$\ln \frac{K_{T_2}}{K_{T_1}} = \frac{E_a}{R} \left( \frac{1}{T_1} - \frac{1}{T_2} \right)$$

$$\text{or } \log \frac{K_{T_2}}{K_{T_1}} = \frac{E_a}{2.303R} \left( \frac{T_2 - T_1}{T_1 T_2} \right)$$

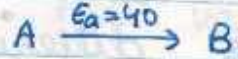
• if  $E_a > 0 \Rightarrow$  on  $\uparrow$  temp.,  $K$  will  $\uparrow$

if  $E_a < 0 \Rightarrow$  on  $\uparrow$  temp.  $K$  will  $\downarrow$

Case I :



$$\frac{K_{410K}}{K_{300K}}$$

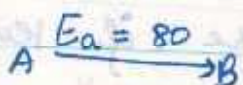


$$\frac{K_{310K}}{K_{300K}}$$

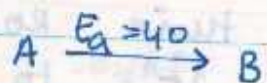
From this data, it can be concluded that % change in value of ' $K$ ' will be more at low temp. as compared to at high temp. for same temp. rise

DR  
Rate of rxn is more sensitive to temp. at low temperature.

Case II :



$$\frac{K_{310K}}{K_{300K}}$$



$$\frac{K'_{310K}}{K'_{300K}}$$

% change will be more where activation energy is high.

### \* Temperature Coefficient ( $\mu$ ) :

It is ratio of rate constant or rate of reaction at 2 temperatures differing by 10 K.

$$\mu = \frac{k_{(T+10)K}}{k_{TK}} = 2-3$$

If temp. is not mentioned then it is taken as 298 K.

Case I : If temp. coeff. is independent of temp.

$$200K \xrightarrow{2} 210K \xrightarrow{2} 220K$$

(actually slightly less than 4 $\times$ )

Q. For rxn A  $\rightarrow$  B temp. coeff. is 3 at  $25^\circ C$ . How many times R.R. will increase if temp. is  $\uparrow$  from  $25^\circ C$  to  $75^\circ C$

$$\rightarrow 3^5 \approx$$

Case II :

If temp. coeff. depends on temp.

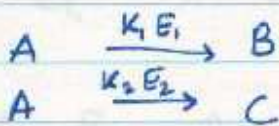
- g. The temp. coeff. ( $\mu$ ) of rxn at 300K is "e". Assuming temp. coeff. to be dependent on temp. what would be the value of

$$\ln \frac{K_{620K}}{K_{300K}} = ?$$

$$\rightarrow \ln \frac{K_{310K}}{K_{300K}} = \frac{E_a}{R} \left( \frac{10}{310 \cdot 300} \right) = 1$$

$$\ln \frac{K_{620K}}{K_{300K}} = \frac{310 \cdot 300}{10} \times \frac{32}{620 \cdot 300} = [16]$$

- g. for 2 parallel rxn



Show that for disappearance of A net activation energy ( $E_3$ ) =  $K_1 E_1 + K_2 E_2$

$$\rightarrow K_1 = A_1 e^{-E_1/RT} \quad \& \quad K_2 = A_2 e^{-E_2/RT}$$

$$K_3 = K_1 + K_2 \Rightarrow A_3 e^{-E_3/RT} = A_1 e^{-E_1/RT} + A_2 e^{-E_2/RT}$$

$\frac{d}{dt} \text{ w.r.t. } T$

$$A_3 e^{-E_3/RT} \left( \frac{E_3}{RT^2} \right) = A_1 e^{-E_1/RT} \left( \frac{E_1}{RT^2} \right) + A_2 e^{-E_2/RT} \left( \frac{E_2}{RT^2} \right)$$

$$K_3 E_3 = K_1 E_1 + K_2 E_2$$

Q. The desorption of gas molecules from the adsorbent obeys Arrhenius eq. The average time upto which a  $N_2$  molecule may remain adsorbed at Pt surface at 400 K is

Given  $A = 1.25 \times 10^8 \text{ s}^{-1}$   
 $E_a = 16 \text{ kcal/mol}$   
 $e^{20} = 5 \times 10^8$ .

$$\rightarrow T_{avg.} = \frac{1}{K_A} = \frac{16 \times 10^3 \times 4.2}{400 \times 8.314} = 14 \text{ s}$$

$$K = 1.25 \times 10^8 e^{-\frac{20}{16}} = \frac{1}{4} = 14 \text{ s} = T_{avg.}$$

Ex 1: 50-53; 56-63, 64, 65-67

\* Differential form of Arrhenius equation:

$$\frac{d(\ln K)}{dT} = \frac{E_a}{RT^2}$$

Case I : if  $E_a$  is independent of temp.

$$\ln \frac{K_2}{K_1} = \frac{E_a}{R} \int_{T_1}^{T_2} \frac{dT}{T^2}$$

$$\Rightarrow \ln \frac{K_{T_2}}{K_{T_1}} = \frac{E_a}{R} \times \frac{(T_2 - T_1)}{T_1 T_2}$$

$$\int d(\ln k) = \frac{E_a}{R} \int \frac{dT}{T^2}$$

$$\Rightarrow \ln k = -\frac{E_a}{R} \times \frac{1}{T} + C$$

$$\Rightarrow \ln \frac{k}{A} = -\frac{E_a}{RT}$$

Case II: if  $E_a = f(T)$

$$E_a = aT^3$$

$$\begin{aligned} & \ln \frac{K_{T_2}}{K_{T_1}} \\ & \int_{\ln K_{T_1}}^{d(\ln k)} = \int_{T_1}^{T_2} \frac{E_a}{RT^2} dT \end{aligned}$$

$$\Rightarrow \ln \frac{K_{T_2}}{K_{T_1}} = \frac{a}{2R} (T_2^2 - T_1^2).$$

Q. find  $E_a$  at 300K if expression of rate constant as fun of temp is given by

$$k = 30 e^{-\frac{40}{T}} \cdot e^{\frac{300}{T^2}}$$

$$\rightarrow k = 30 e^{\frac{300-40T}{T^2}}$$

$$-\frac{E_a}{R} = \frac{--}{T^2}$$

$$E_a = - \left( \frac{300-40T}{T} \right) RT = \underline{\underline{38 R}} \quad \times$$

$$\ln k = \ln 30 - \frac{40}{T} + \frac{300}{T^2}$$

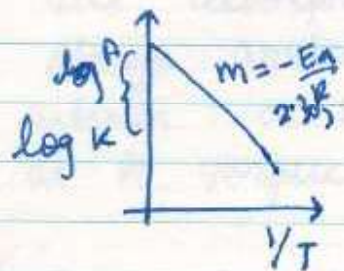
$$\Rightarrow E_a = R \left( 40 - \frac{600}{T} \right)$$

$$\Rightarrow \frac{d(\ln k)}{dT} = 0 + \frac{40}{T^2} - \frac{600}{T^3}$$

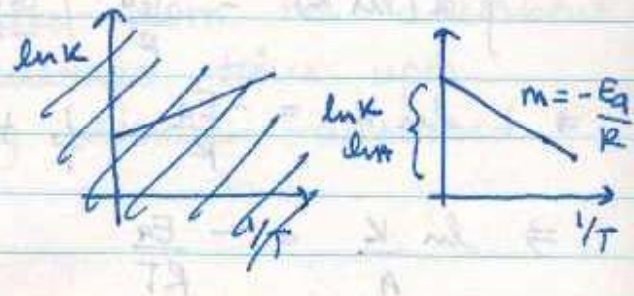
$$= 38 \text{ R}$$

## \* Graphical Representation

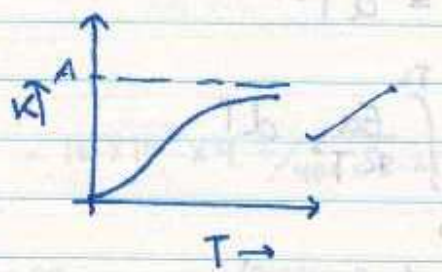
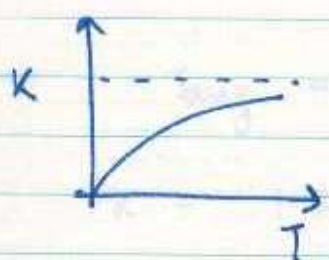
$$1. \log K \text{ v/s } 1/T$$



$$2. \ln K \text{ v/s } \frac{1}{T}$$



$$3. K \text{ v/s } T$$



$$K = A e^{-E_a/RT}$$

After rearranging the terms in the above eqn  
we get  $\ln K = \ln A - \frac{E_a}{R} \left( \frac{1}{T} \right)$

$\therefore \ln K = \text{constant} - \frac{E_a}{R} \left( \frac{1}{T} \right)$

$$\frac{\partial \ln K}{\partial T} = -\frac{E_a}{R} \left( \frac{1}{T^2} \right) = -\frac{E_a}{RT^2}$$

## \* Theories of Chemical Rxn :

### 1) Collision Theory :

- This theory was given by Arrhenius & Wentorf
- This theory is applicable mainly for bimolecular or gases rxn.

According to it, a chemical rxn takes place due to collision b/w reactant molecules coz during collision old bonds are broken and new bonds are formed.

Under normal conditions, collision frequency ( $Z_{11}$ ) is equal to  $10^{30}$  ~~most~~ collisions / s.l

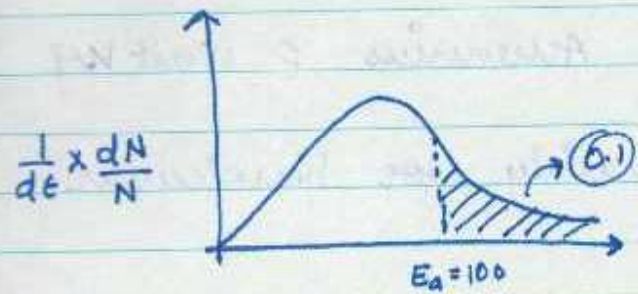
If all these collisions were effective then every rxn would have been completed within a fraction of time but this is not the case. It means all collisions are not effective.

• Effective collision :> These are the collisions which cause product formation

In order to be effective collision, it should satisfy following two criteria :

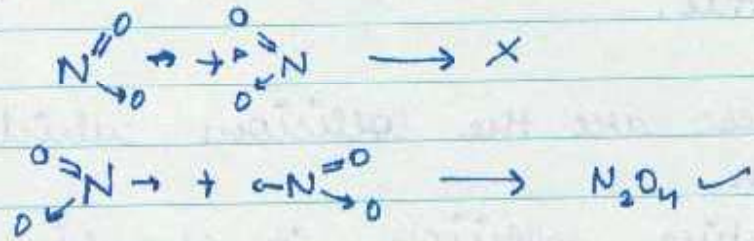
- i) Energy barrier
- ii) Orientation factor

- Energy Barriers : Colliding molecules must have sufficient K.E. so that it can cross Activation / Threshold barrier.



Fraction of molecules having sufficient KE =  $e^{-E_a/RT}$

- Orientational factor : All colliding molecules having sufficient KE should also have proper orientation in order to get converted into product.



- Steric factor (P) : It represents the fraction of molecules having proper orientation.

Considering all above factors, rate of chemical reaction can be given by

RDR = collision freq.  $\times$  effective collision

= collision freq.  $\times$  collision with sufficient energy  $\times$  proper orientation

$$= \left( \pi (\tau_A + \tau_B)^2 u_{avg} N_A^* N_B^* \right) \times e^{-E_a/RT} \times P$$

$$= \left( \pi (\tau_A + \tau_B)^2 u_{avg} \times N_0^2 \times P \right) e^{-E_a/RT} \times [A][B]$$

$$\text{Rate} = A \cdot e^{-E_a/RT} [A][B]$$

$$= k[A][B]$$

$$K = A \cdot e^{-E_a/RT} f$$

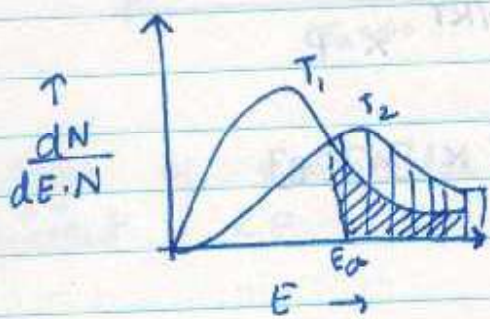
$$A = \pi (\tau_A + \tau_B)^2 u_{avg} \times N_0^2 \times P$$

$$A \propto u_{avg} \propto \sqrt{T}$$

Actually 'A' depends on temp. but on small change in temp., its value does not change significantly that is why it is considered as independent of temp.

- Why on  $10^{\circ}\text{C}$  rise in temperature R.R. becomes double?

→ On  $\uparrow$  temp., fraction of molecules having sufficient K.E. becomes increased drastically. As a result rate becomes double or more than that.

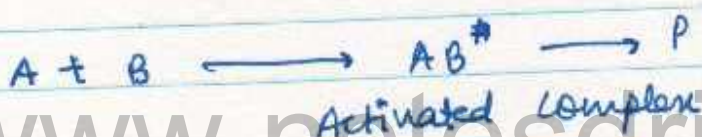


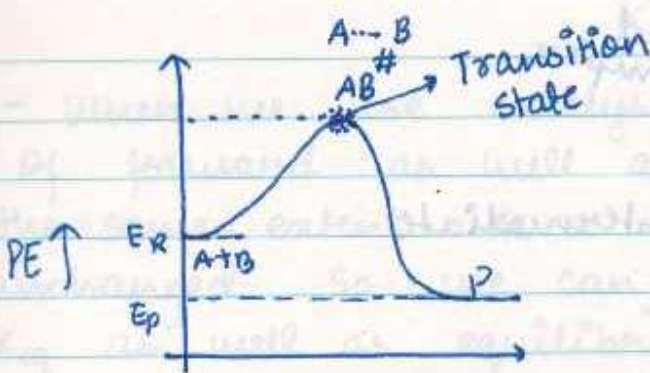
### \* Activated Complex Theory:

- ACT formerly known as transition state theory offers following advantage over GT.:

- i) It can be applied for gaseous reactants as well as for the rans taking place in liquid soln.
- ii) Steric factor appears automatically in contrast to collision theory where 'P' has to be grafted.

### Reaction Profile:

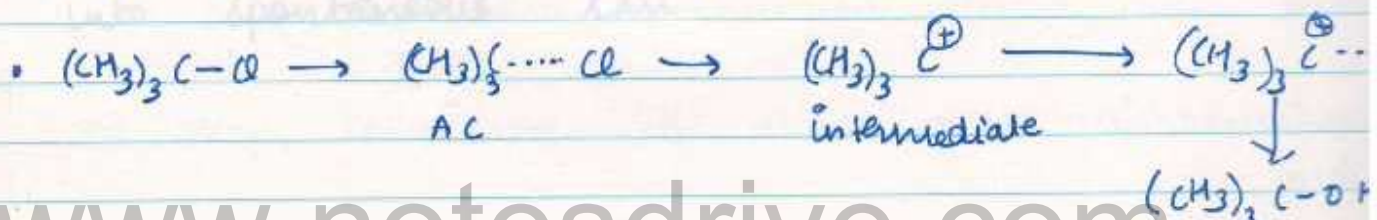
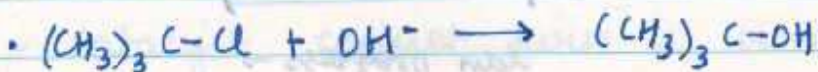


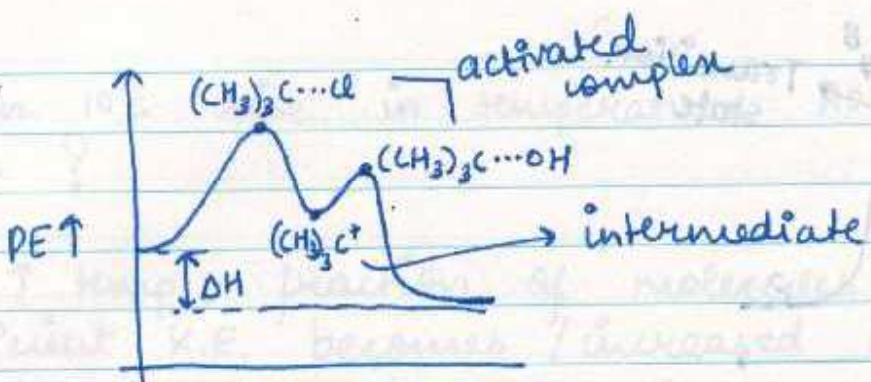


- Initially only reactants A & B are present.
- As rxn proceeds, A & B come into contact, disintegrate & begin to exchange atoms.
- As P.E.  $\uparrow$  to a maximum, the cluster of atoms corresponds to the region near maxima are referred as activated complex. The config<sup>n</sup> at actual peak of curve is called transition state.
- After maxima P.E. falls as atoms rearrange to form product.
- In fact all  $AB^{\ddagger}$  will not proceed to form product. Some of  $AB^{\ddagger}$  will decay back to reactants.

Note :  $AB^{\ddagger}$  is not an intermediate compd in rxn but rather it is a structure that is in process of change in direc<sup>n</sup> of product.

SN<sub>1</sub>,





### \* CATALYST :

- These are the substances which ↑ the rate of rxn without actually being consumed.
  - Catalyst participates qualitatively not quantitatively.
  - Catalyst increases the rate of chemical rxn by providing an alternative path with lower activation energy.
  - Catalysed rxns are generally complex rxns
- ii
- uncatalysed

catalysed

PET

$\downarrow \Delta H$

Rxn progress →

N.C.E.M

uncatalysed

catalysed

PET

Rxn progress →

- When we use catalyst it increases the rate of forward as well as backward rxn to the same extent. As a result,  $K_{eq}$  remains unchanged. So we can say, it does not alter  $K_{eq}$  as well as equilibrium concentrations or partial pressures of rxn components.

For uncatalysed,

$$K_{eq} = \frac{k_f}{k_b} = \frac{A_f e^{-E_{af}/RT}}{A_b e^{-E_{ab}/RT}} = A e^{-(E_{af}-E_{ab})/RT}$$

For catalysed,

$$k_f' = A_f e^{-(E_{af}-x)/RT} = A_f e^{-E_{af}RT} \cdot e^{x/RT}$$

$$k_b' = A_b e^{-(E_{ab}-x)/RT} = A_b e^{-E_{ab}/RT} \cdot e^{x/RT}$$

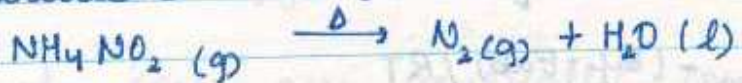
$$K_{eq} = \frac{k_f'}{k_b'} = A e^{-(E_{af}-E_{ab})/RT}$$

- Catalyst does not alter value of  $\Delta H$  or  $\Delta G$  for rxn.
- Catalyst cannot turn a non-spontaneous rxn into spontaneous rxn.

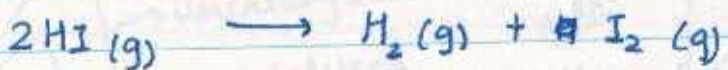
- \* MECHANISM OF RXN:
- It represents a path through which the reactants get converted into products.
- During mechanism rxn is explained step wise.
- Rate determining step (RDS): This is the slowest step of rxn. (If rxn takes place in more than one step).
- Molecularity: The no. of reacting species (atoms, ions, molecules) taking part in an elementary rxn which must collide simultaneously in order to bring about a chemical change.

Types:

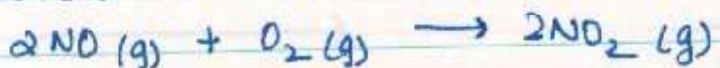
- a) Unimolecular Rxn:



- b) Bimolecular Rxn:



- c) Trinolecular Rxn:



- Some imp. points regarding molecularity:

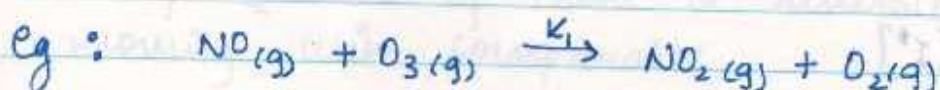
- \* Molecularity can never be fractional, -ve or zero.

- \* Probability of simultaneous collision of three or more molecules is very low. As a result runs with molecularity 4 or more are not observed.
  - \* In case of elementary ran, molecularity as well as order will be same.
  - \* There is no meaning of molecularity for complex ran, while order can be defined for complex ran. However, molecularity can be defined for each step of a complex ran coz each step will be an elementary ran.
- Q. An elementary ran can't be of fractional order.
- True.

Case I : If reactants directly get converted to product.

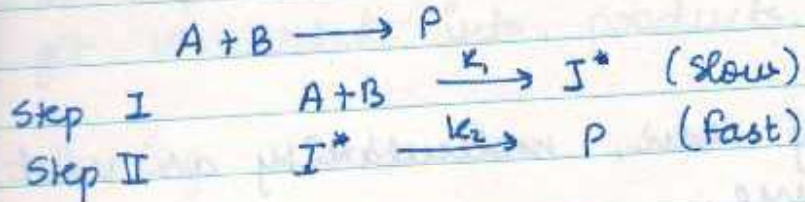


$$\text{Rate} = k_1 [A][B]$$

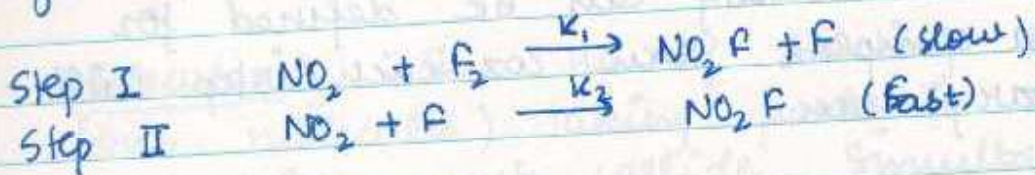
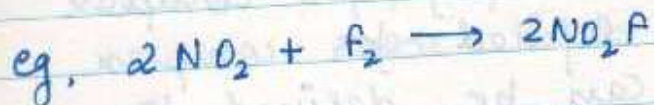


$$\text{Rate} = k_1 [NO_2][O_3]$$

Case II : If rxn takes place in more than one step & one of them is slow

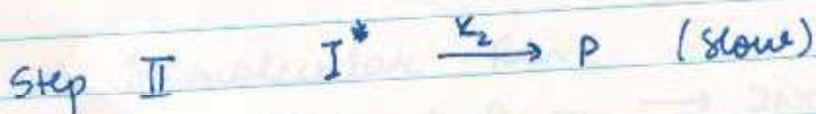
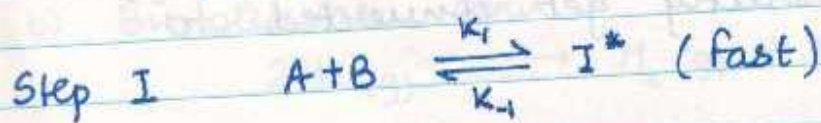


$$\text{Rate} = k_1 [A][B]$$



$$\text{Rate} = k_1 [\text{NO}_2][\text{F}_2]$$

Case III : If rxn takes place in more than one step & one of them is slow, & one is reversible.



$$\text{Rate} = k_2 [I^*]$$

$$= k_2 \times \frac{k_1}{k_{-1}} \times [A][B]$$

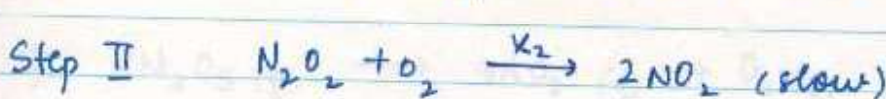
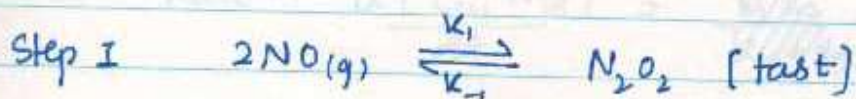
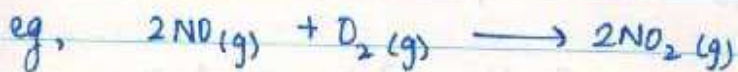
$$ROR = K_{0x} [A][B]$$

$$K_{0x} = \frac{k_1 k_2}{k_{-1}} = \frac{A_1 e^{-E_1/RT} \cdot A_2 e^{-E_2/RT}}{A_{-1} \cdot e^{-E_{-1}/RT}}$$

$$K_{0x} = \frac{A_1 A_2}{A_{-1}} \times e^{-(E_1 + E_2 - E_{-1})/RT}$$

$$\text{Overall} = E_1 + E_2 - E_{-1}$$

Activation Energy (overall) can be +ve, -ve or zero.

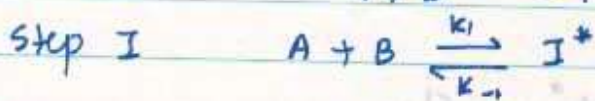
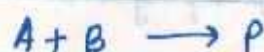


$$\frac{k_1}{k_{-1}} = \frac{[\text{N}_2\text{O}_2]}{[\text{NO}]^2}$$

$$ROR = k_2 [\text{N}_2\text{O}_2] [\text{O}_2]$$

$$= k_2 \cdot \frac{k_1}{k_{-1}} \cdot [\text{NO}]^2 [\text{O}_2]$$

Case IV : If rxn takes place in more than one step & one of them is reversible & steps are moving with comparable.



$$\text{Rate} = \frac{d[P]}{dt} = k_2 [I^*]$$

We can apply Steady state on  $[I^*]$

$$\frac{d[I]}{dt} = 0$$

$$\text{Rate} = \frac{k_1 k_2}{k_1 + k_2} [A][B]$$

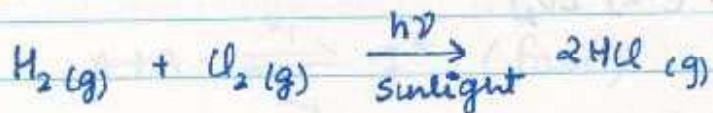
If  $k_1 \gg k_{-1} \gg k_2$ ,

$$\text{Rate} = \frac{k_1 k_2}{k_{-1}} [A][B]$$

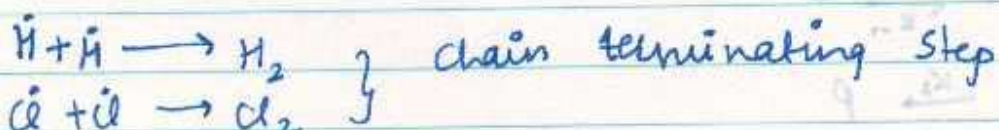
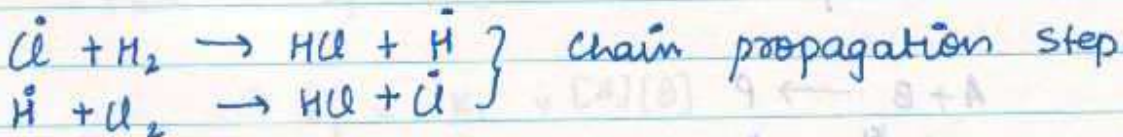
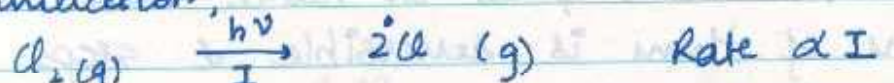
If  $k_2 \gg k_{-1}$ ,

$$\text{Rate} = k_1 [A][B]$$

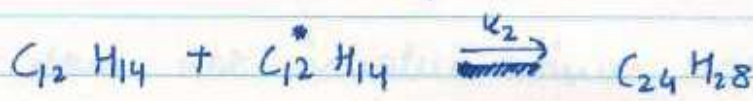
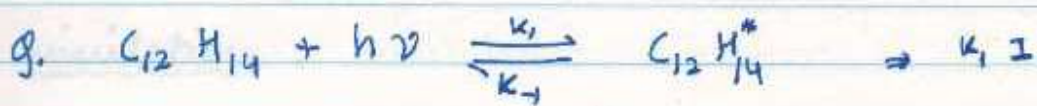
\* Photochemical Rxn:



Chain initiation,



- first step of photochemical rxn will be of zero order. Its rate will depend on intensity of light ( $I$ ).
- Photochemical rxn will be complex.
- Photochemical rxn can have any order.

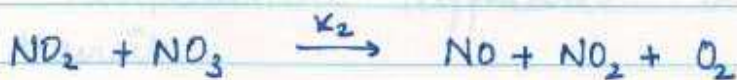
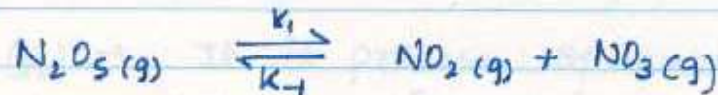


$$\begin{aligned} k_1 - k_2[C_{12}H_{14}] \\ - k_2[C_{12}H_{14}^*] = 0 \end{aligned}$$

$$\rightarrow \text{Rate} = \frac{d[C_{24}H_{28}]}{dt} = \frac{k_1 k_2}{k_1 + k_2} [C_{12}H_{14}] \cdot [k_1 - k_2[C_{12}H_{14}]]$$



Mech



$$\text{P.T. } \frac{d[O_2]}{dt} = k[N_2O_5]$$