# **Chemical Kinetics**

## Rate of Reaction:

The quantity of a reactant species consumed or the quantity of a product species formed in unit time in a chemical reaction is called the rate of the reaction. Consider a reaction,  $A \rightarrow B$ 

$$Rate = -\frac{d[A]}{dt} = +\frac{d[B]}{dt}$$

-ve sign indicates that with progress of time concentration of A will decrease.

### Rate Laws & Rate Constant:

Let us consider a general reaction,

$$A + B \rightarrow Product$$

According to rate law,

$$Rate \propto [A][B]$$

$$Or, Rate = K[A][B]$$

This is known as rate law & 'K' is called rate constant which is independent of concentration but depend on temperature.

Mention the factors affect the rate of a chemical reaction.

Factors affecting the rate of a chemical reaction are following -

- i. Nature of reactants
- ii. Concentration of reactants
- iii. Effect of temperature
- iv. Surface area of reactants
- v. Presence of catalyst
- vi. Presence of light

### Order of a Reaction:

Order of a reaction is defined as the sum of the powers or exponents to which the molar concentrations are raised in the rate law or rate equation.

Let us consider a general reaction,

 $lA + mB \rightarrow Product$ 

According to rate law,

#### $R = K[A]^a[B]^b$

where a & b may or may not be equal to l & m respectively. Here the overall order of the reaction is (a + b).

- $\checkmark$  Order of a reaction is purely experimental quantity.
- ✓ Order of reaction may be fraction, zero or even negative. As for example, thermal decomposition of acetaldehyde,

 $CH_3CHO \rightarrow CH_4 + CO$ 

has order equal to 3/2.

### Example of Reactions having Different Order:

- I. <u>Zero order reaction</u>: Dissociation of ammonia on tungsten or dissociation of *HI* on gold surface.
- II. <u>First order reaction</u>: Conversion of cyclopropane to propylene, substitution reaction of tertiary alkyl halide by nucleophile.
- III. Second order reaction:  $H_2 + I_2 \rightarrow 2HI$ , substitution reaction of primary alkyl halide by nucleophile.
- IV. <u>Fractional order reaction</u>: Thermal decomposition of acetaldehyde,

$$CH_3CHO \rightarrow CH_4 + CO$$

has order equal to  $\frac{3}{2}$ .

### \*<u>Elementary Reaction:</u>

A chemical reaction that takes place in one & only one step, i.e., all that occurs in a single step is called elementary reaction. For example –

$$PCl_5 \rightleftharpoons PCl_3 + Cl_2$$
$$2HI \rightleftharpoons H_2 + I_2$$

Molecularity of an elementary reaction is the number of the reactant particles that are involved in each individual chemical event.

### Molecularity of Elementary Reaction:

It is defined as the number of reactant particles (atoms, molecules, free radicals or ions) that are involved in each individual chemical event.

If a reaction takes place in a number of steps then slowest step is usually the rate determining step. Molecularity of the rate determining step is said to be the molecularity of the reaction.

- ✓ Molecularity of a reaction is purely theoretical concept.
- ✓ Molecularity of a reaction is necessarily a positive integer.

Elementary reactions with molecularity greater than three generally do not occur – justify.

The reactants are activated as a result of molecular collisions. But the collisions generally require high amount of energy because of inter – nuclear & inter – electronic repulsions. For a collision minimum two molecules are must. If the molecularity of an elementary step is 3, then three molecules must collide requiring high amount of energy. For a four – body collision, energy required will be still higher & therefore unknown. It is far easier to have two – body collisions & that will take place easily. Therefore, elementary reaction of molecularity 4 or greater is unknown. If molecularity becomes greater than 3, then such reaction involve two or more steps, such that each step has its own molecularity not greater than 3.

### Difference Between Molecularity & Order:

The main differences are –

- I. Order is an experimental quantity whereas molecularity of a reaction is a purely theoretical concept.
- II. Order of a reaction is not necessarily integer. It may be fraction, zero or even negative. Whereas molecularity of a reaction is necessarily a positive integer.
- III. Order may change with physical condition, such as pressure, solvent etc whereas molecularity is an invariant number for a given chemical reaction.

Give an example of a chemical reaction where molecularity & order is different.

 $CH_3CHO \rightleftharpoons CH_4 + CO$ 

So, rate =  $k[CH_3CHO]^{3/2}$ . Here the order of the eaction is 1.5 but molecularity is 1. Hence order & molecularity is different.

#### Integrated Rate Laws:

 Zero Order Reaction: A reaction whose rate is independent of concentration is called a zero order reaction. Thus for a zero order reaction –

$$A \rightarrow pdt.$$
  
So, rate =  $-\frac{d[A]}{dt} = k[A]^0 = k$ 

where [A] is the concentration of the reactant A at time t, thus -

$$\int_{a}^{[A[} d[A] = -k \int_{0}^{t} dt$$
$$Or, [A] - a = -kt$$
$$Or, [A] = a - kt$$

A zero order reaction will be completed when [A] = 0, so,

$$a - kt_{completion} = 0$$
  
Or,  $t_{completion} = \left(\frac{a}{k}\right)$ 

<u>Graphical Representation</u>: If we plot [A] vs t, we get a straight line with slope = -k.



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<u>Half – life period</u>: When  $t = t_{\underline{1}}$ , then, [A] = a/2, now –

$$\frac{a}{2} = a - kt_{\frac{1}{2}}$$
  
Or,  $t_{\frac{1}{2}} = \left(\frac{a}{2k}\right)$ , *i.e.*,  $t_{1/2} \propto a$ 

Unit of rate constant: mol. L<sup>-1</sup>sec<sup>-1</sup>

Characteristics:

- I. Rate is independent of instantaneous concentration of the reactant.
- II. Unit of rate & rate constant are the same.
- III. Zero order reaction must be multistep in nature.
- IV. Zero order reaction can be completed in a finite time.
- V. Half life period is directly proportional to initial concentration of reactant.

Write down the unit of the rate constant of a zero order reaction.

#### $mol. L^{-1}sec^{-1}$

What do you mean by half-life period of a reaction?

The half-life of a reaction,  $t_{1/2}$ , is the amount of time needed for a reactant concentration to decrease by half compared to its initial concentration.

Zero order reaction is multistep — Explain.

If zero order reaction is single step then the reaction would be an elementary reaction & in that case order of the reaction would be same as molecularity. But molecularity of a reaction cannot be zero. So zero order reaction must be a multistep reaction. The reactant with respect to which the order becomes zero is involved in a step which is not the rate determining step (r.d.s.).

Does a zero order reaction go to completion?

Rate equation for a zero order reaction —

$$-\frac{dc}{dt} = kC^{0}$$
  

$$Or, \int dc = -\int kdt$$
  

$$Or, C = -kt + IC$$
  

$$at t = 0, C = C_{0}, then IC = C_{0}$$
  

$$So, C = C_{0} - kt$$

Hence, a zero order reaction will go to completion & time duration is  $\frac{C_0}{\nu}$ .

> For a reaction the plot of log  $t_{1/2}$  versus log a turns out to be a straight line with positive slope and intercept, which makes an angle of  $45^{\circ}$  with the log a axis. What is the order of the reaction?

We know half - life period of a nth order reaction is given as -

$$t_{1/2} = \frac{k}{a^{n-1}}$$
  
Or, log  $t_{1/2} = \log k + (1-n) \log a$ 

Slope of  $\log t_{1/2} vs \log a = (1 - n)$ . So we can write –

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 $(1-n) = tan 45^0 = 1, \quad Or, \quad n = 0$ 

Hence the given reaction is a zero order reaction.

In a certain kinetic study, the plot of  $\log t_{1/2} vs \log a$  was found to be a straight line with unit slope. State with reason the order of the reaction.

We know half - life period of a nth order reaction is given as -

$$t_{1/2} = \frac{\kappa}{a^{n-1}}$$
  
Or, log  $t_{1/2} = \log k + (1-n) \log a$ 

Slope of  $\log t_{1/2}$   $vs \log a = (1 - n)$ . So we can write –

$$(1-n) = 1$$
$$Or, n = 0$$

Hence the given reaction is a zero order reaction.

 First Order Reaction: A first order reaction is one whose rate is linearly proportional to the instantaneous concentration of the reactant. Let us consider a first order reaction –

$$A \rightarrow B$$

Let initial concentration of reactant is  $[A]_0 = a \& at an arbitrary time t$  it is [A] = a - x. Then the differential rate law is given as –

$$-\frac{d(a-x)}{dt} = k(a-x)$$
$$Or, \left(\frac{dx}{dt}\right) = k(a-x)$$
$$Or, \int_{0}^{x} \frac{dx}{(a-x)} = k \int_{0}^{t} dt$$
$$Or, \ln\left(\frac{a}{a-x}\right) = kt$$
$$Or, \log\left(\frac{a}{a-x}\right) = \frac{k}{2.303}t$$
$$Or, t = \frac{2.303}{k} \log\left(\frac{a}{a-x}\right)$$

<u>Graphical Representation</u>: If we plot  $\log (a_a - x) vs t$ , we get a straight line passing through the origin & slope = k/2.303.



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<u>Half – life period</u>: When  $t = t_{\frac{1}{2}}$ , then,  $x = \frac{a}{2}$ , from above equation –

$$t_{\frac{1}{2}} = \frac{2.303}{k} \log\left(\frac{a}{a - \frac{a}{2}}\right) = \frac{1}{k} \ln 2 = \frac{0.693}{k}$$

So, half – life period is independent of initial concentration of reactant. Unit of rate constant:  $sec^{-1}$ 

Characteristics:

- I. First order reaction never goes to completion.
- II. Half life period is independent of initial concentration of reactant.

First order reaction never proceeds to completion – justify or criticize. For first order reaction –

$$\frac{[A]}{[A]_0} = e^{-kt}$$

When reaction will be completed then [A] should be equal to zero & it will be possible only when  $t = \infty$ , so first order reaction never goes to completion.

Show for a first order reaction, the time required for 99.9% completion of the reaction is almost 10 times the time for 50% completion.

For a first order reaction,

$$t = \frac{1}{k} ln \frac{a}{a-x} = \frac{2.303}{k} log \frac{a}{a-0.999a}$$
  

$$Or, t = \frac{2.303}{k} log 10^3 = \frac{6.909}{k}$$
  

$$We \ know, t_{\frac{1}{2}} = \frac{0.693}{k}$$
  

$$So, \frac{t}{t_{\frac{1}{2}}} = \frac{6.909}{0.693} \approx 10.$$
  

$$Hence, t = 10 \times t_{1/2} \ (proved)$$

Show that the time required for 75% completion of a first – order reaction is twice the time required for 50% completion of the reaction.

For a first order reaction,

$$t = \frac{1}{k} ln \frac{a}{a-x} = \frac{2.303}{k} log \frac{a}{a-0.75a}$$
  

$$Or, t = \frac{2.303}{k} log 4 = \frac{1.3865}{k}$$
  

$$We \ know, t_{\frac{1}{2}} = \frac{0.693}{k}$$
  

$$So, \frac{t}{t_{\frac{1}{2}}} = \frac{1.3865}{0.693} \approx 2.$$
  
Hence,  $t = 2 \times t_{1/2}$  (proved)

3. <u>Second Order Reaction</u>: Consider a second order reaction involving only one reactant, i.e.

$$A \rightarrow pdt$$
.

Let initial concentration of reactant is  $[A]_0 = a$  & at an arbitrary time *t* it is [A] = a - x. Then the differential rate law is given as –

$$-\frac{d(a-x)}{dt} = k(a-x)^{2}$$
$$Or, \left(\frac{dx}{dt}\right) = k(a-x)^{2}$$
$$Or, \int_{0}^{x} \frac{dx}{(a-x)^{2}} = k \int_{0}^{t} k \, dt$$
$$Or, \left(\frac{1}{a-x}\right) - \frac{1}{a} = kt$$
$$Or, \frac{1}{a-x} = \frac{1}{a} + kt$$
$$Or, kt = \frac{x}{a(a-x)}$$
$$Or, t = \frac{1}{k} \times \frac{x}{a(a-x)}$$

<u>Graphical Representation</u>: If we plot 1/[R] *vs t*, we get a straight line with intercept = 1/a & *slope* = k.



<u>Half – life period</u>: When  $t = t_{\frac{1}{2}}$ , then,  $x = \frac{a}{2}$ , from above equation – <u>Half – life period</u>: When  $t = t_{\frac{1}{2}}$ , then,  $x = \frac{a}{2}$ , from above equation –

$$t_{\frac{1}{2}} = \frac{1}{k} \times \frac{\frac{a}{2}}{a \cdot \frac{a}{2}} = \frac{1}{ak}$$

Hence, half – life period is inversely proportional to initial concentration of reactant. <u>Unit of rate constant</u>:  $mol^{-1}lit.sec^{-1}$ 

Characteristics:

- I. Second order reaction transfers to first order reaction when one of the reactant is present in large excess.
- II. Half life period of second order reaction is inversely proportional to initial concentration of reactant.

#### Chemical Kinetics (VJRC)

Write down the unit of rate constant of a 2<sup>nd</sup> order reaction.

$$mol^{-1}lit.sec^{-1}$$

Second order reaction never proceeds to completion – justify or criticize.

For a 2nd order reaction,  $k = \frac{1}{t} \frac{x}{a(a-x)}$ 

So, 
$$t = \frac{1}{k} \frac{x}{a(a-x)}$$
  
When  $t = t_{completion}$ , then  $a = 0$ ,  
So,  $t_{completion} = \frac{1}{k} \cdot \frac{x}{0} = \infty$ 

So, 2<sup>nd</sup> order reaction never goes to completion.

For a second order reaction A —> products. Hence show that the time required for three – fourths of the reactant to decay (t<sub>3/4</sub>) is equal to 3t<sub>1/2</sub>.

For a second order reaction —

$$\frac{1}{a-x} = \frac{1}{a} + kt$$

$$Or, \frac{1}{a-\frac{3a}{4}} = \frac{1}{a} + kt_{\frac{3}{4}} (for\frac{3}{4} th of the reactant)$$

$$Or, \frac{1}{a} = \frac{1}{a} + kt_{\frac{3}{4}}$$

$$Or, kt_{\frac{3}{4}} = \frac{4}{a} - \frac{1}{a} = \frac{3}{a}$$

$$Or, t_{\frac{3}{4}} = 3 \times \frac{1}{ka} = 3t_{1/2} (we know for 2nd order reaction t_{\frac{1}{2}} = \frac{1}{ka})$$

$$Or, t_{\frac{3}{4}} = 3 \times t_{1/2} (proved)$$