

- RATE OF A CHEMICAL REACTION Time taken OR Increase in conc of reactant
- AVERAGE RATE
- Consider a reaction: A + B C+ D
- $-\frac{\Delta[A]}{\Delta t} = -\frac{\Delta[B]}{\Delta t} = \frac{\Delta[C]}{\Delta t} = \frac{\Delta[D]}{\Delta t}$
- INSTANTANEOUS RATE
- Consider a reaction: aA + bB cC + dD $-\frac{1}{\alpha}\frac{d[A]}{dt} = -\frac{1}{b}\frac{d[B]}{dt} = \frac{1}{c}\frac{d[C]}{dt} = \frac{1}{d}\frac{d[D]}{dt}$
- Unit of Rate = mol litre<sup>-1</sup> s<sup>-1</sup>
- QDuring the decomposition of  $H_2O_2$ , 48 g  $O_2$ is formed per minute at a certain point of time. The rate of formation of water at this point is
- (a) 0.75 mol min <sup>-1</sup> (b) 1.5 mol min<sup>-1</sup>
- (c) 2.25 mol min<sup>-1</sup> (d) 3.0 mol min<sup>-1</sup>

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### FACTORS INFLUENCING **RATE OF REACTION**

Factors	Effect on reaction rate
e in concentration	Increases
e in temperature	Increases
e of catalyst	Increases
	Factors e in concentration e in temperature se of catalyst

QWhich of the following will lead to an increase in the rate of the reaction?

- a) Decrease in temperature
- b) Decreasing concentration of reactants
- c) Addition of catalyst
- d) Addition of inhibitor

### RATE LAW

Consider a general reaction, aA + bB → product

Rate = k[A]° [B]b (law of mass action)

- Rate =  $k[A]^x [B]^y$  (rate law expression)
- x & y are determined experimentally and may or may not be equa
- x & y represents the order of reaction with respect to A & B

### RATE CONSTANT

- Larger the value of k, faster is the reaction.
- The value of k changes only with temperature for given reaction.

 $(unit of rate constant = (mol)^{1-n} L^{n-1} s^{-1}$ 

- QThe rate constant of a zero-order reactions has the unit
- (a) s <sup>-1</sup>

- (b) mol L-1 s-1
- (c) L<sup>2</sup> mol<sup>-2</sup> s<sup>-1</sup>
- (d) L mol<sup>-1</sup> s<sup>-1</sup>

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### ORDER AND MOLECULARITY

Consider a general reaction,

aA + bB → product

Rate =  $k[A]^x [B]^y$ 

molecularity = a + border = x + y

Molecularity

An experimentally

It can be equal to zero positive, negative and fraction It cannot be zero, fracti infinite and imaginary.

Order

- Q When the rate of the reaction is equal to the rate constant, the order of the reaction is
- (a) zero order

- (b) first order
- (c) second order
- (d) third order

### PSEUDO ORDER REACTIONS

Consider the reaction

 $C_{12}H_{22}O_{11} + H_2O \xrightarrow{H^+} C_{\Lambda}H_{12}O_{\Lambda} + C_{\Lambda}H_{12}O_{\Lambda}$ 

In these reactions, concentration of water (one of the reactants) is in excess and its concentration remains constant throughout

Thus, rate < [ $C_{12}H_{22}O_{11}$ ]

Therefore, order= 1

Q For a pseudo first-order reaction, what is the unit of the rate of the reaction?

- (a) s<sup>-1</sup>
- (b) mol L<sup>-1</sup>s <sup>-1</sup>
- (c) mol<sup>-1</sup> L s<sup>-1</sup>
- (d) mol<sup>-2</sup> L<sup>2</sup> s<sup>-1</sup>

# CHEMICAL KINETICS

### ELEMENTARY & COMPLEX REACTIONS

- Reactions occurring only in one step are called elementary reactions while that involving a sequence of elementary reactions, are called complex reactions. ■ In case of complex reactions, the slowest step is called rate
- determining step. From slow sten

Note Consider the reaction 20,-30, Step - 1  $O_3 \rightleftharpoons O_2 + O \text{ (fast)}$ 

 $r = k [O] [O_a]$  $[O] \propto [Q_3]$ ; From fast step  $r = k [O_3]^2 [O_2]^{-1}$  $\frac{\text{Step - 2}}{\text{O + O}_3} \rightarrow 2\text{O}_2 \text{ (slow)}$ 

### **INTEGRATED RATE EQUATIONS** Zero order

 $k = \frac{[A]_{0} - [A]}{t}$ First order

 $k = \frac{2.303 \log [A]_0}{1.00}$ 

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

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## HALF LIFE PERIOD

Zero order  $t_{\frac{1}{2}} = \frac{[A]_0}{2k}$ First order

 $t_{\frac{1}{2}} = \frac{0.693}{1}$ Second order

# FIRST ORDER TRICKS $t_{75\%} = 2t_{\frac{1}{2}}$

 $t_{\frac{1}{2}} = \frac{1}{k [A]_0}$ 

# $t_{90\%} = \underline{2.303}$ $t_{99.9\%} = 10t_{\frac{1}{2}}$

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### GRAPHICAL REPRESENTATION

**ARRHENIUS EQUATION** 

 $k = Ae^{-E\alpha/RT}$  $\log k = \log A - \frac{E_a}{2.303R} \left( \frac{1}{T} \right)$ 

> ${}^{ullet}$  For every  $10^{\circ}$  rise in temperature, rate becomes double and hence, rate constant becomes double.

A reaction with higher value of E will have smaller value of rate constant.

## QSuppose the reaction: $A + 2B \rightarrow AB_2$ occurs by the following mechanism:

Step 1 :  $A + B \rightarrow AB$  slow Step 2 :  $AB + B \rightarrow AB_2$  fast

Overall A + 2B  $\rightarrow$  AB<sub>2</sub>

(a) k[A] (b) k[B] (c) k[A][B] (d)  $k[B]^2$ 

### Q A first order reaction has a specific reaction rate of $10^{-2} \text{ sec}^{-1}$ . How much time will it take for 20 g of the reactant to reduce to 5 g?

- (a) 138.6 sec (b) 346.5 sec
- (c) 693.0 sec (d) 238.6 sec

## Q The half-life period of zero order reaction is directly proportional to the

- a) Rate constant
- b) Initial concentration of reactants
- c) Final concentration of reactants
- d) Concentration of products

### O The graph of t<sub>1</sub> versus initial concentration 'a' is for

- a) First order
- b) Second order
- c) Zero order
- d) Can't predict

# Q The slope of Arrhenius plot (In k vs $\frac{1}{T}$ ) of first order reaction is - $5 \times 10^3$ K. The value of E of the reaction is [Given: R=8.314 J K<sup>-1</sup> mol<sup>-1</sup>)

(a)  $-83 \text{ kJ mol}^{-1}$  (b)  $41.5 \text{ kJ mol}^{-1}$ 

(c) 83 kJ mol <sup>-1</sup>

(d) 166 kJ mol <sup>-1</sup>