

MANZIL

For JEE Aspirants

Legend



#ATDB.uno
GOC



Rohit Agrawal (RA Sir)

TOPICS TO BE COVERED



1. Electronic Displacement Effect



2. Acid and Base



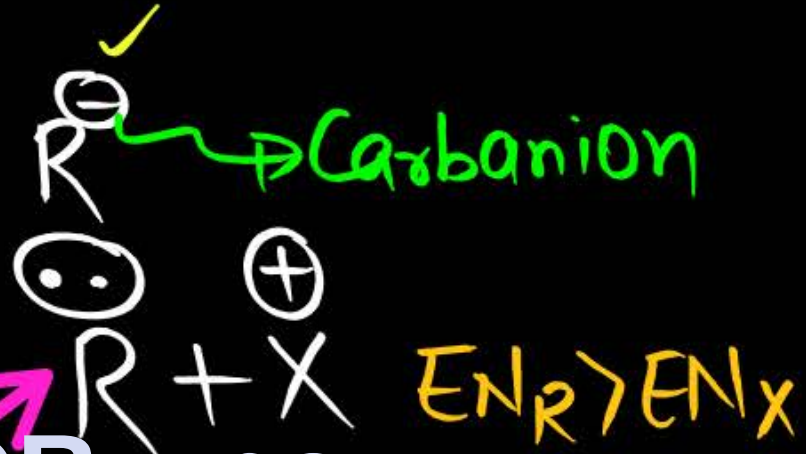
ATDB.uno



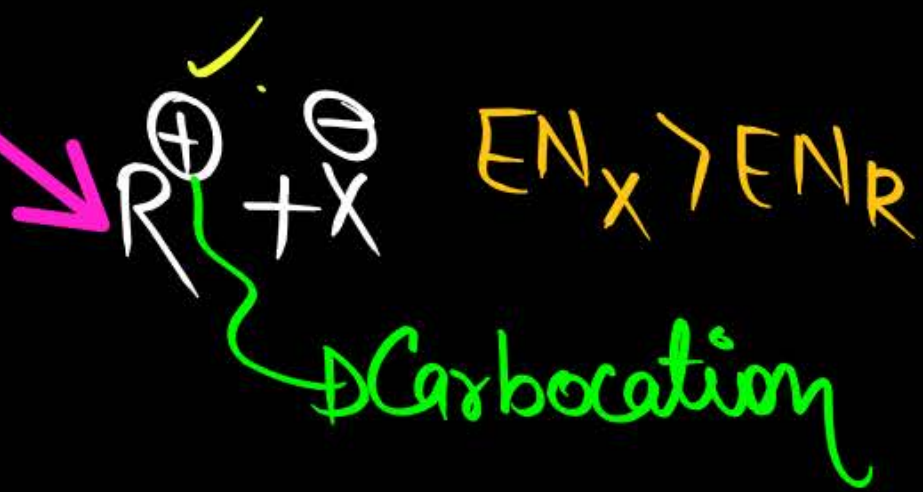
Bond fission



Homolysis
Heterolysis

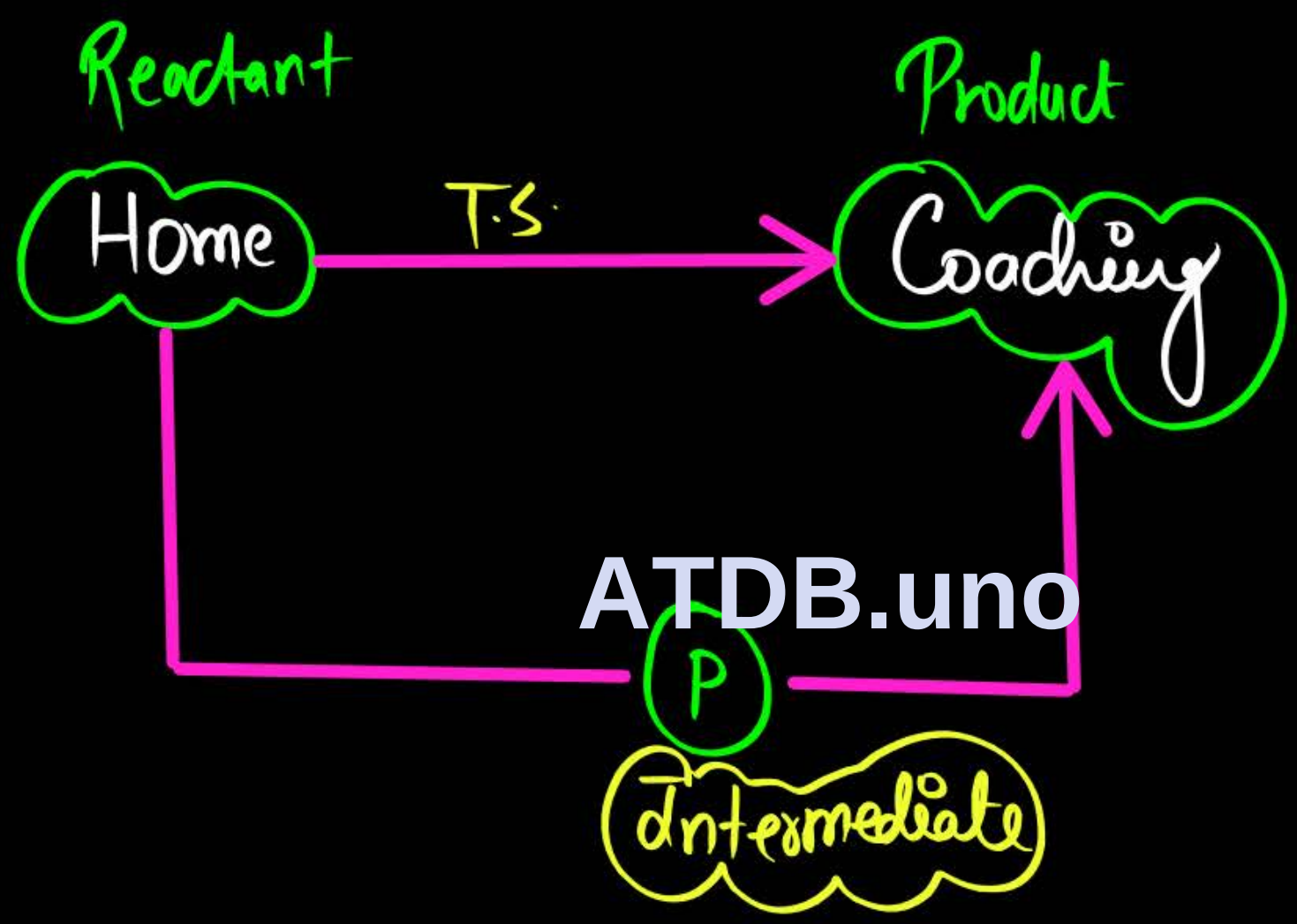


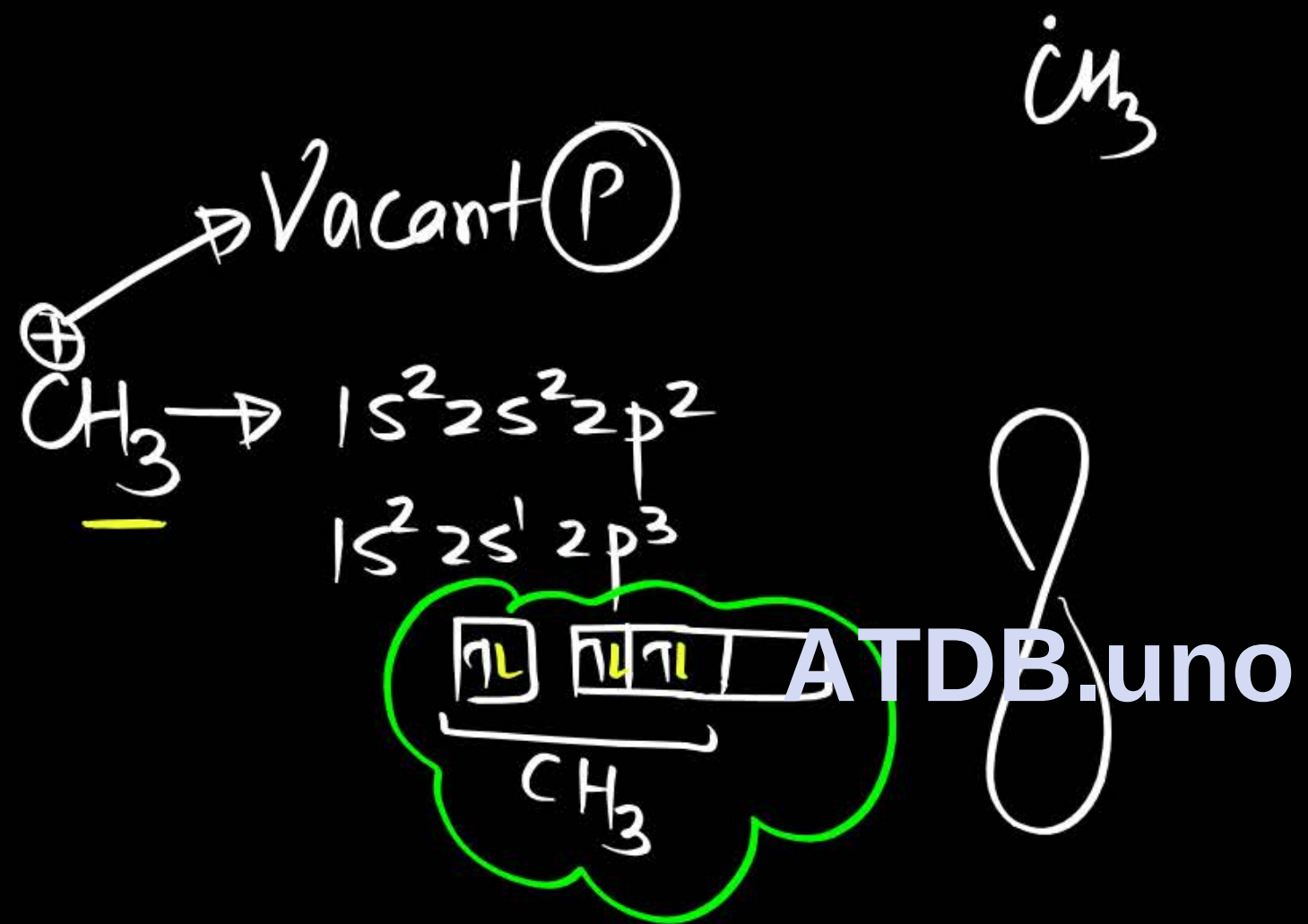
ATDB.uno

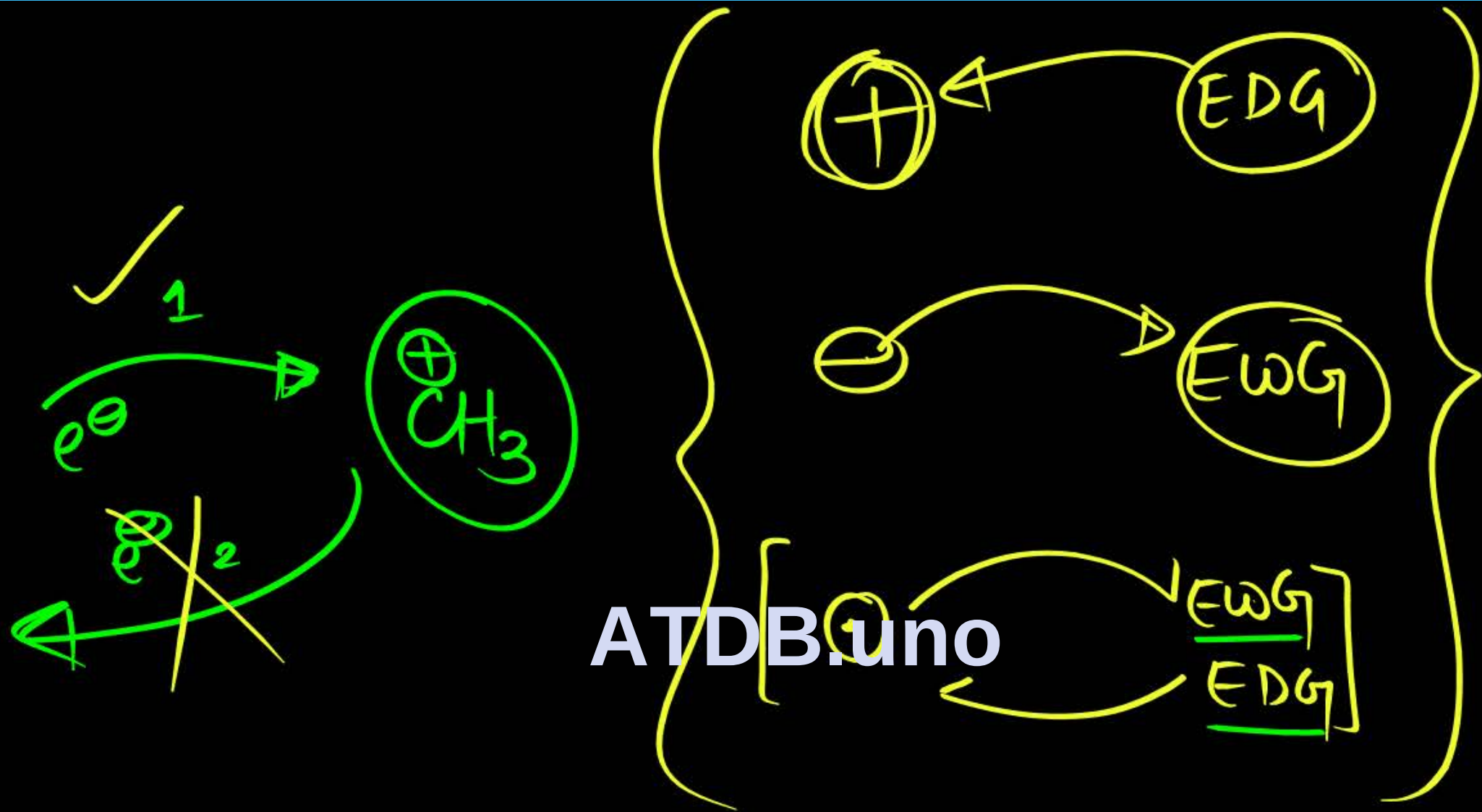




Intermediate







ATDB.uno



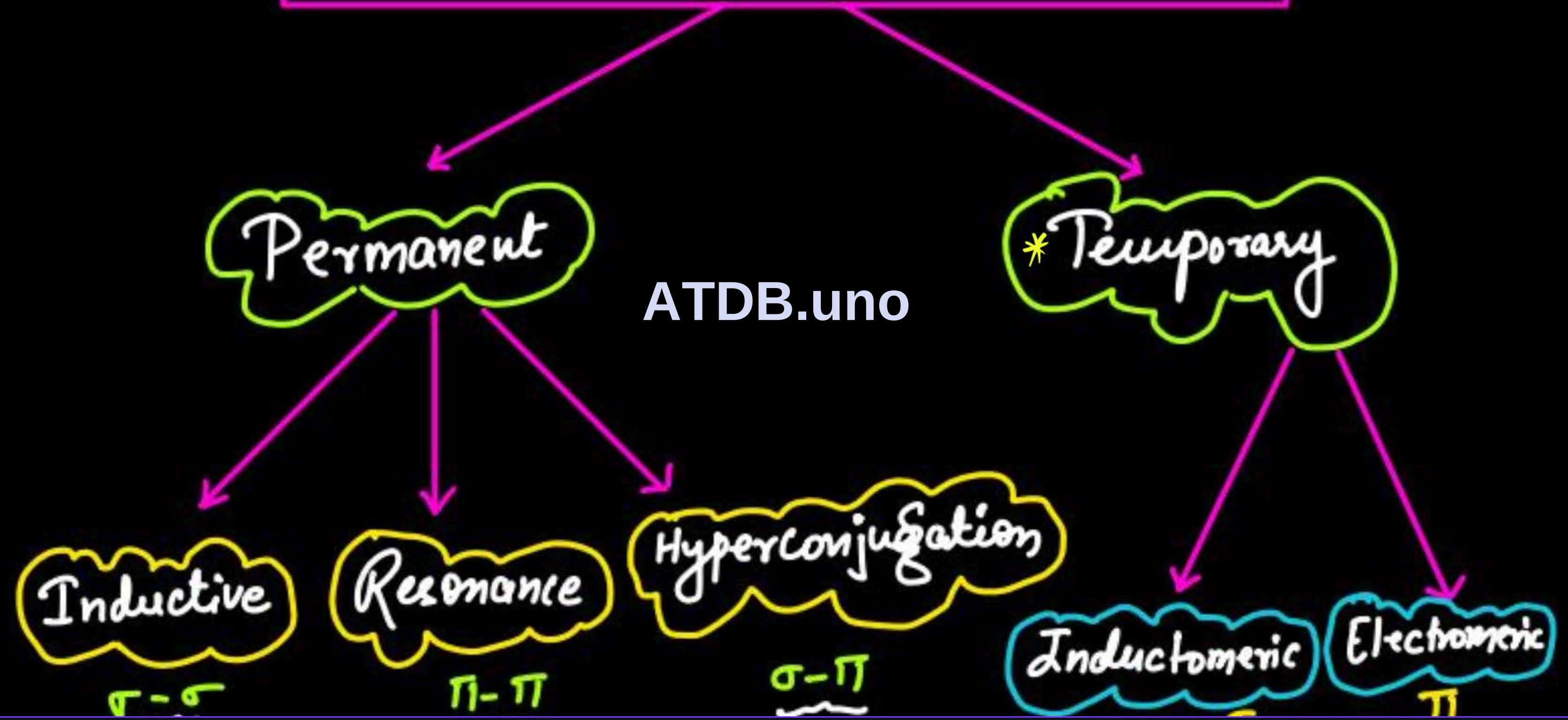


✓✓ Electronic Displacement Effect

- The electron displacements due to the influence of an atom or a substituent group present in the molecule cause permanent polarisation of the bond.

ATDB.uno

Electronic Displacement Effect

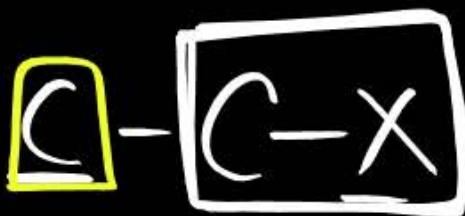




Weakest effed

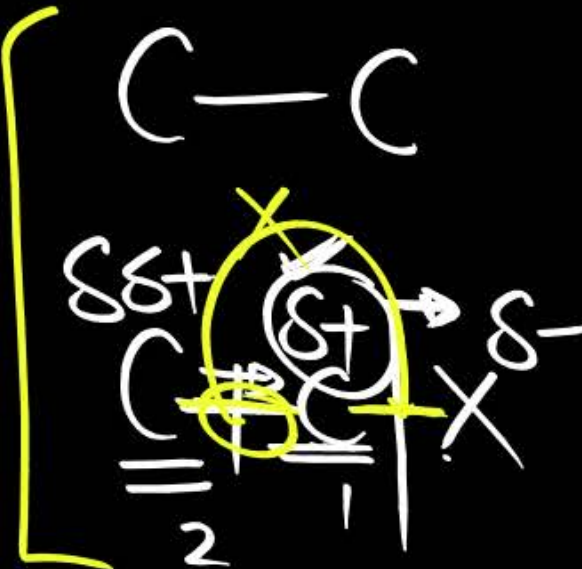


* Inductive effect



✓ EN

ATDB.uno



* e⁻ shifting



Inductive Effect

- When a covalent bond is formed between atoms of different electro negativity, the electron density is more towards the more electronegative atom of the bond. Such a shift of electron density results in a polar covalent bond.
- This shifting of electron density is known as ‘Induction’.
- Due to this induction, adjacent Atom gets effected and developed polarity. This is known as “Inductive Effect”

Properties Of Inductive Effect



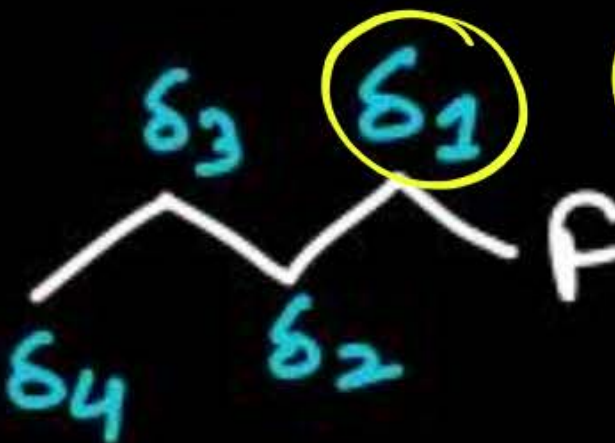
- It is permanent and partial displacement of sigma electrons.
- It is a permanent effect and weak effect.
- Electrons will never changes its atomic orbital.
- Inductive effect is an additive effect.
- It is a distance dependent effect as distance increases inductive effect decreases.
- Only 10% of original charge transfers in each bond and $\delta_4 \approx 0$.



Find the order of charges ?



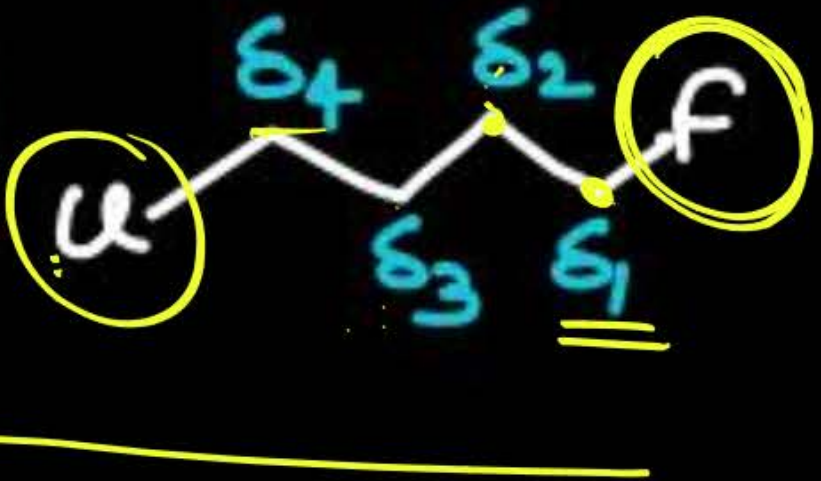
(1)



1 7 2 7 3 7 4

ATDB.uno

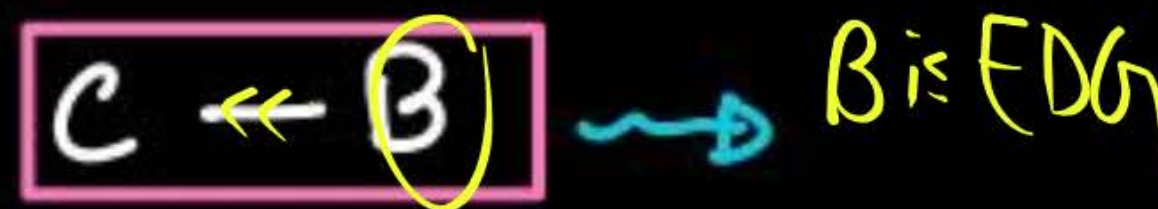
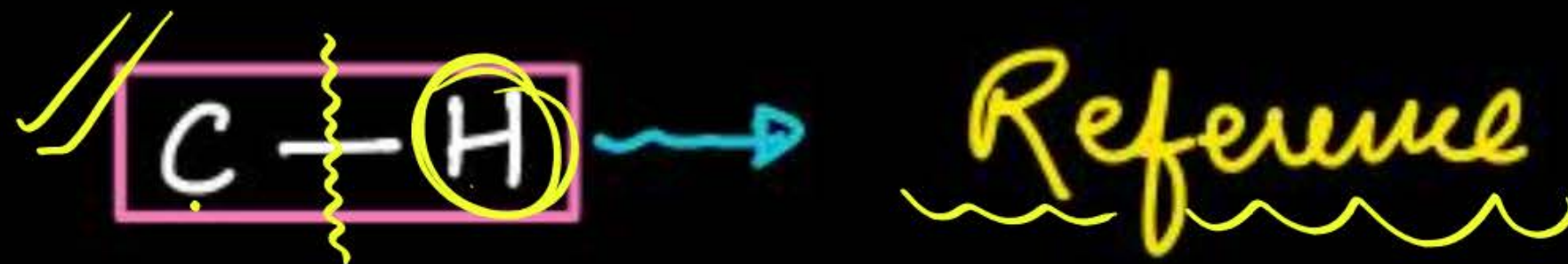
(2)



1 7 4 7 2 7 3

Types Of Inductive Effect

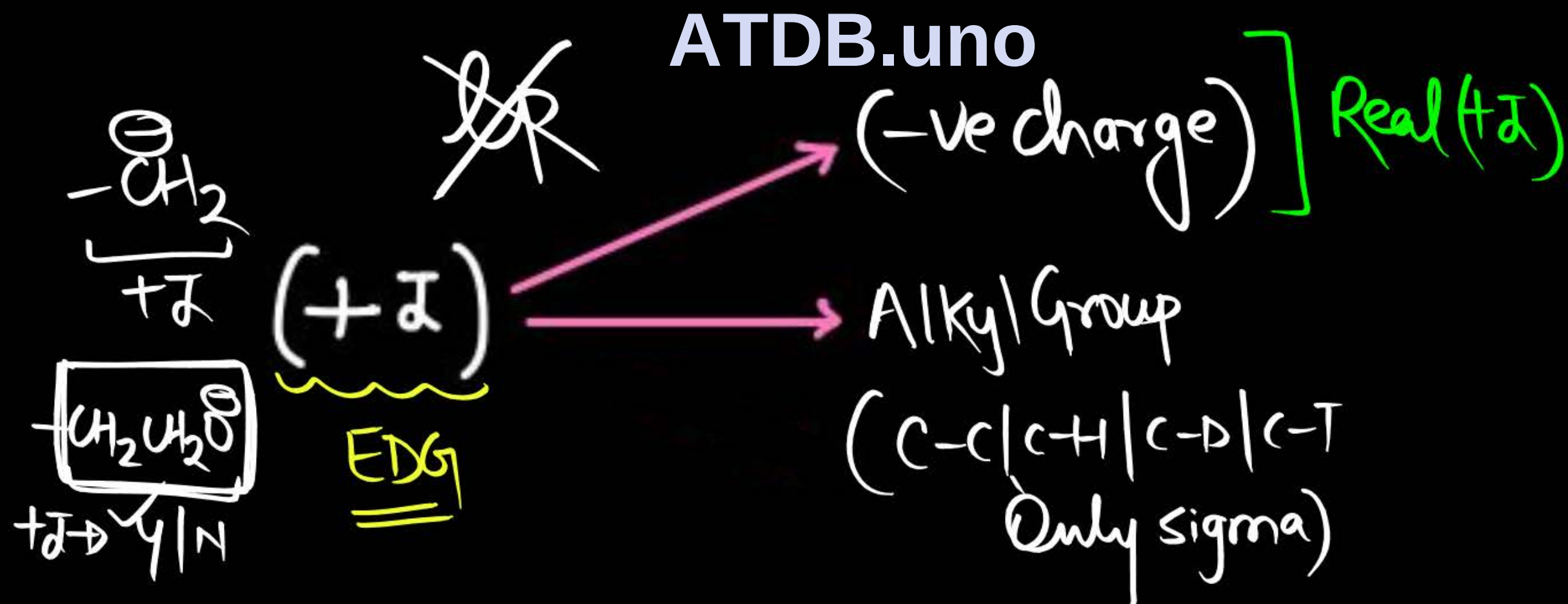
EDG|EWG

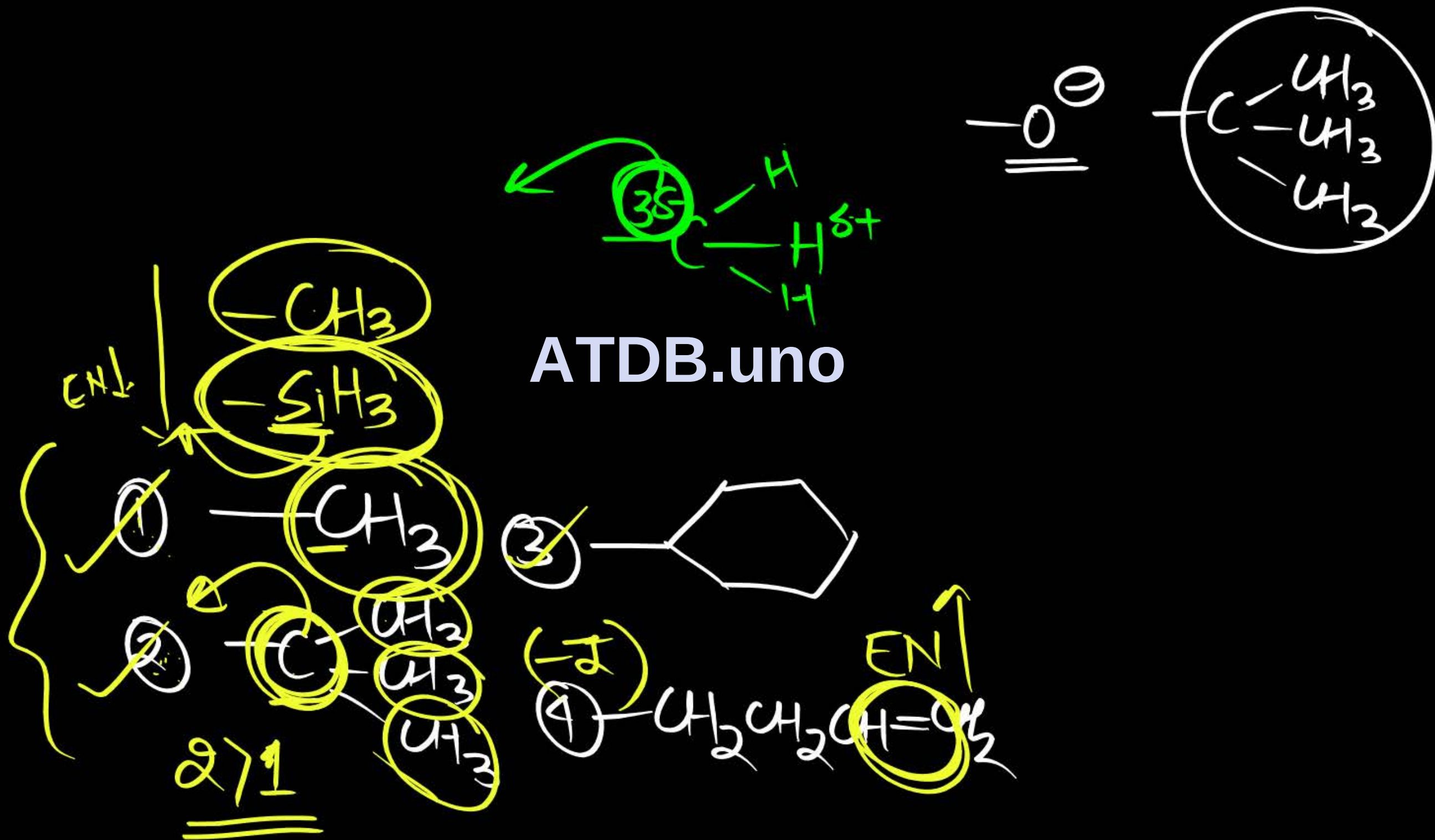


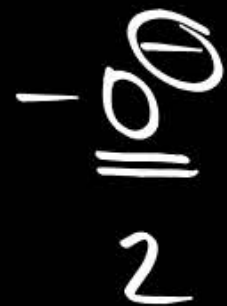
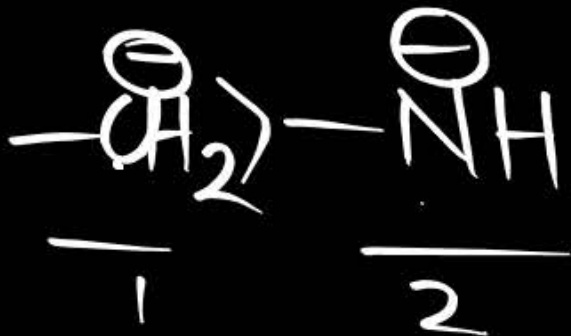
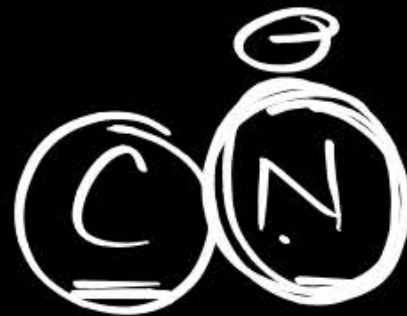


Electron Donating Effect (+I)

- When bonded pair of electrons are shifted away from any atom or group w.r.t. C—H bond then atom or group shows electron donating effect.



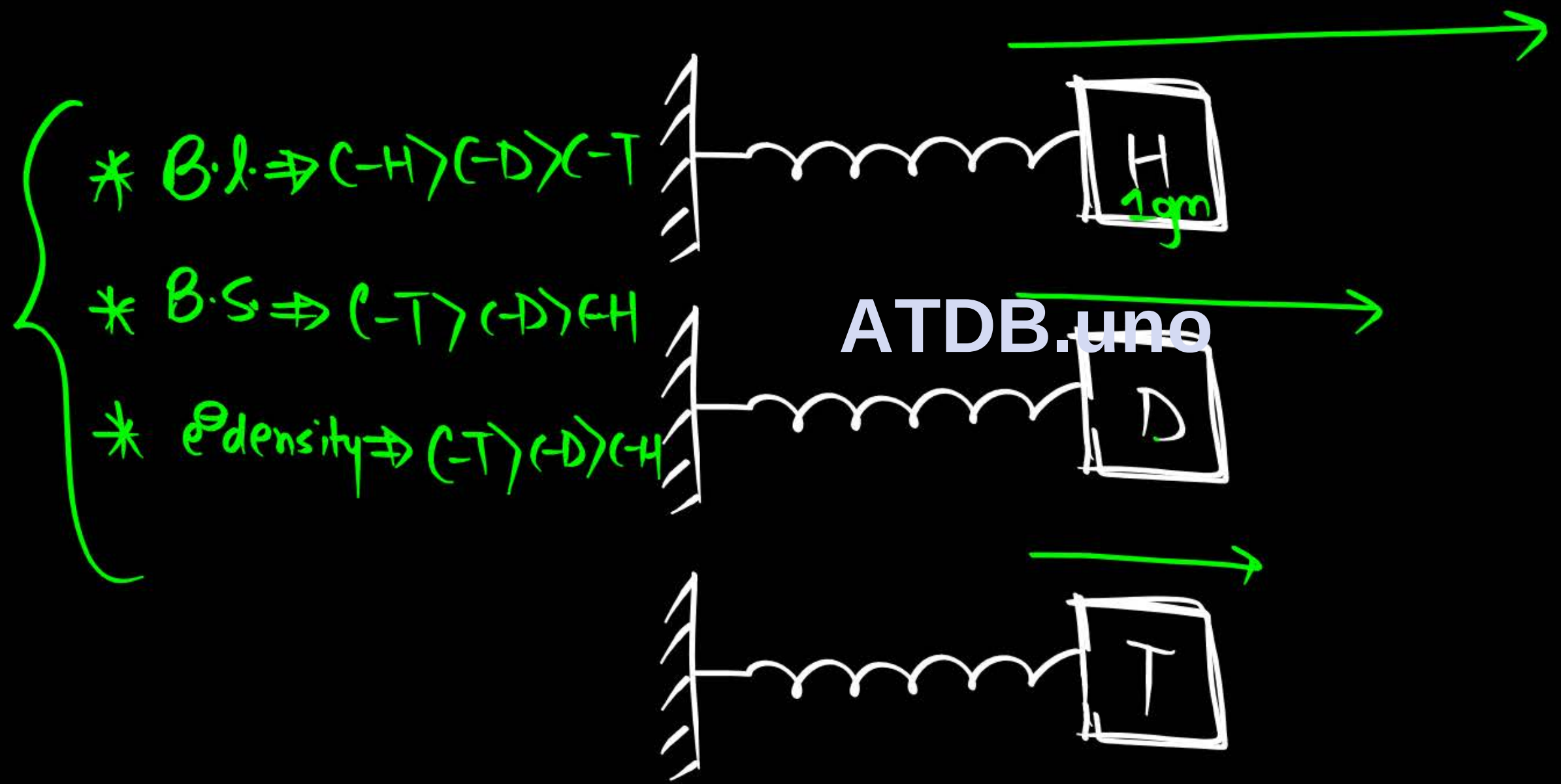




ATDB.uno



ATDB.uno

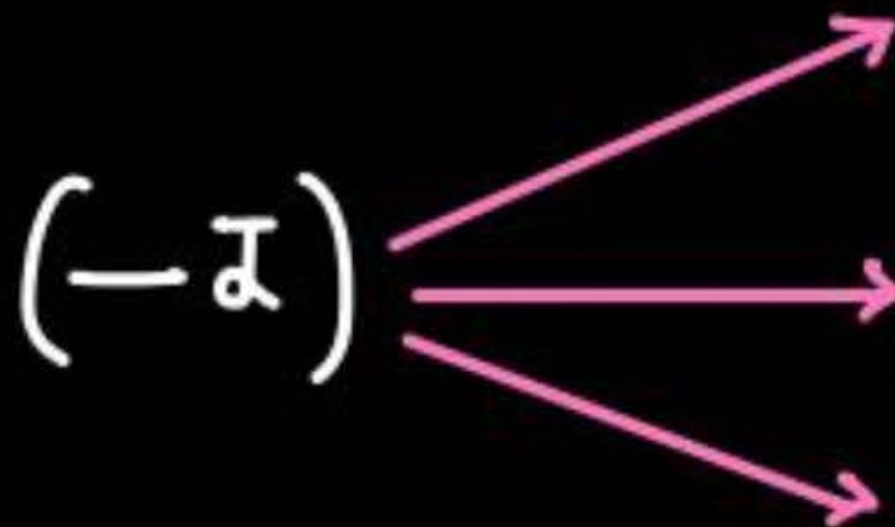




Electron Withdrawing Effect (—I)

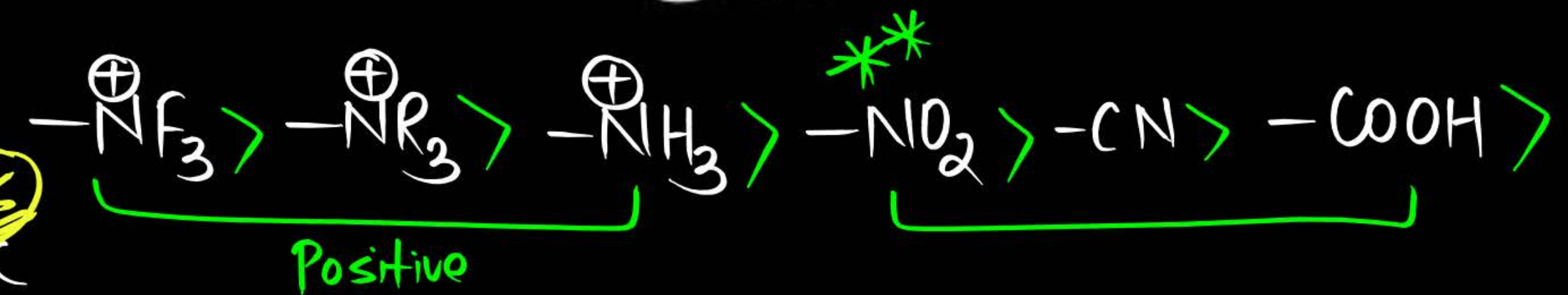
- When bonded pair of electrons are shifted towards any atom or group w.r.t. C—H bond then atom or group shows electron withdrawing effect.

ATDB.uno

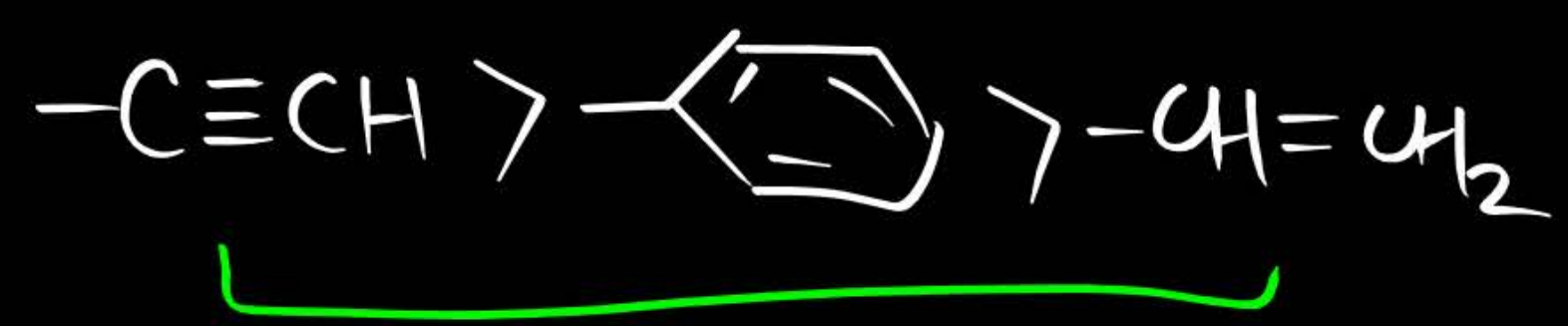




-I Series



3.25
C=C
2.75
1 C=C
3 C=C





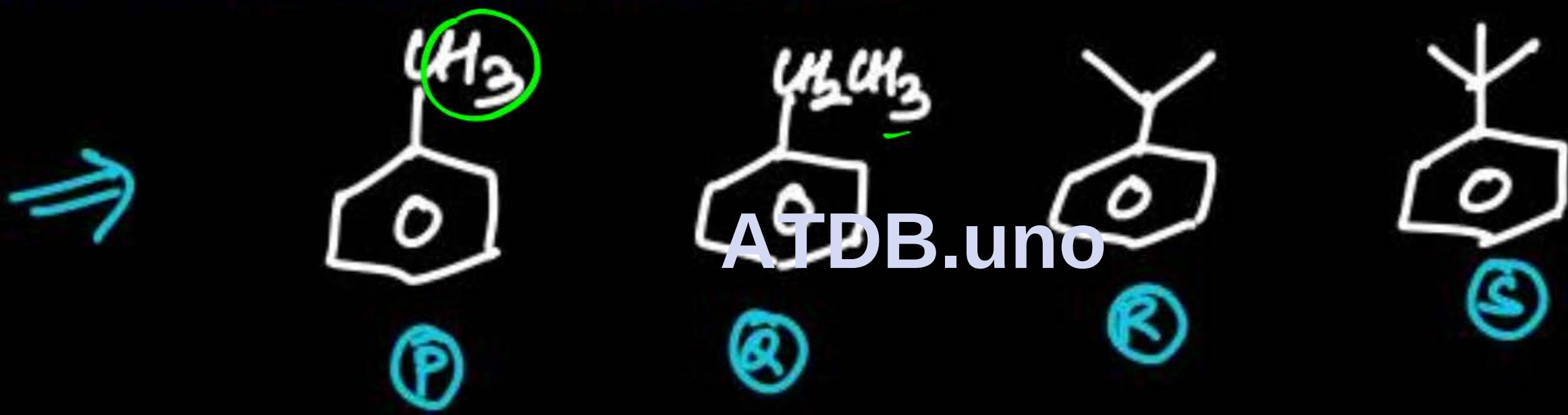
E^+ density

Nathan & Baker

ATDB.uno

Hyperconjugation

Experiment- Baker and Nathan

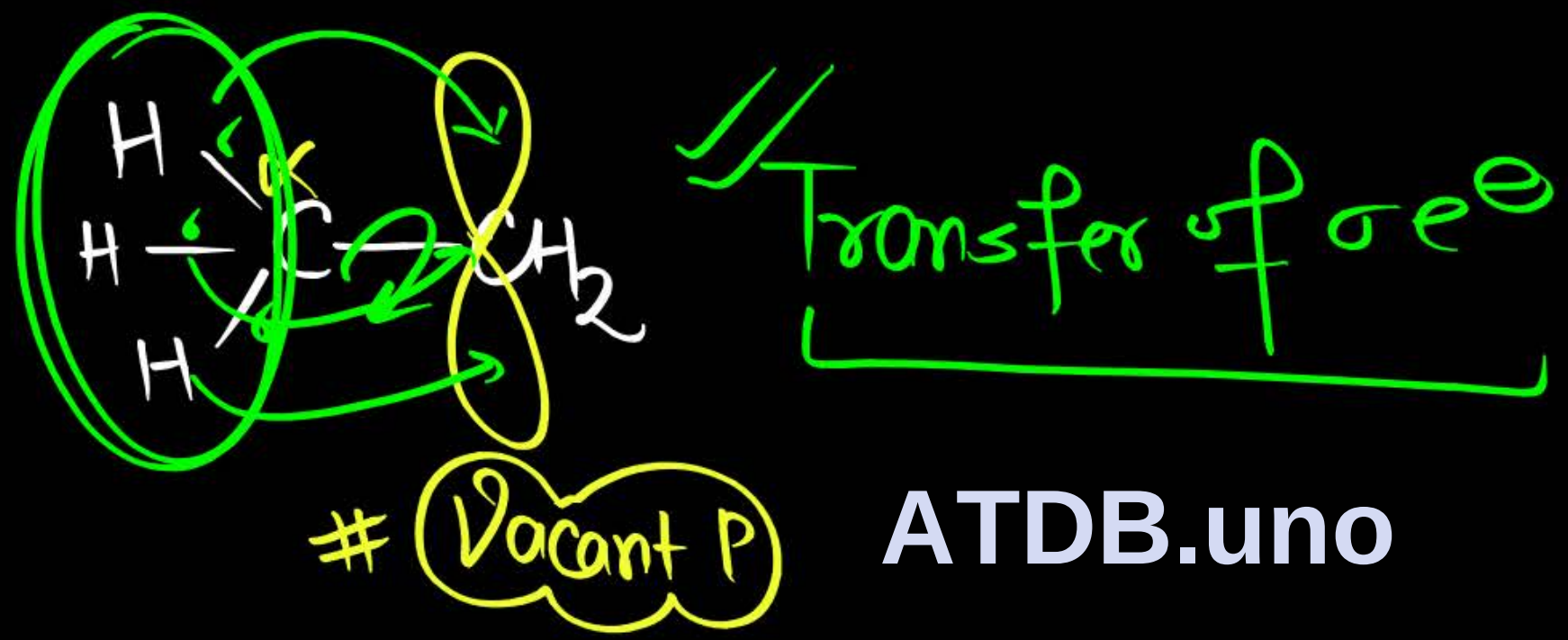


Rate of E^+ ⇒

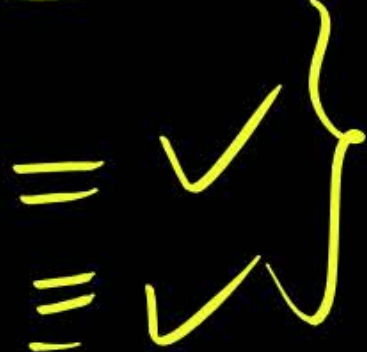
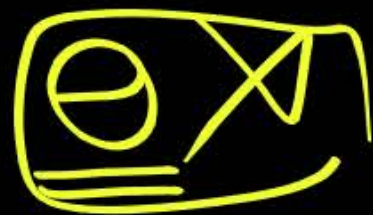
S > R > Q > P (expected)

P > Q > R > S (Observed)

Also known as:
Baker-Nathan effect



ATDB.uno

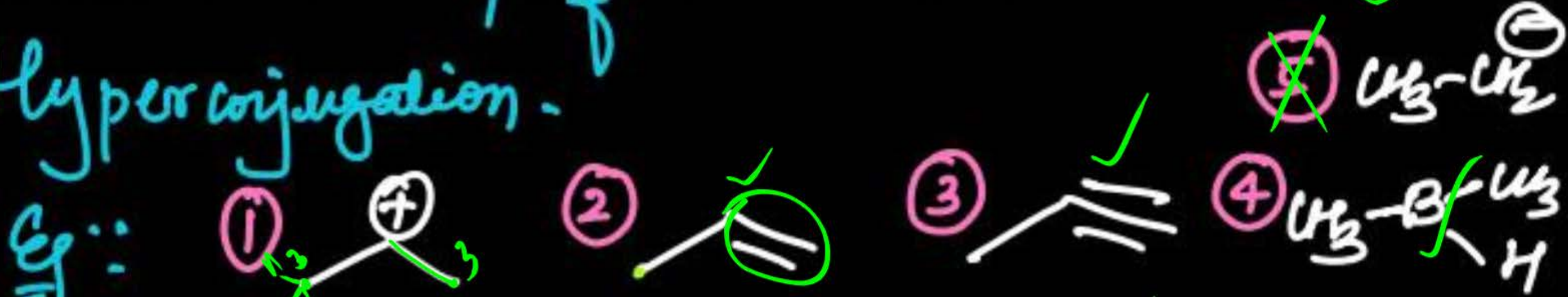


Hyperconjugating Structure

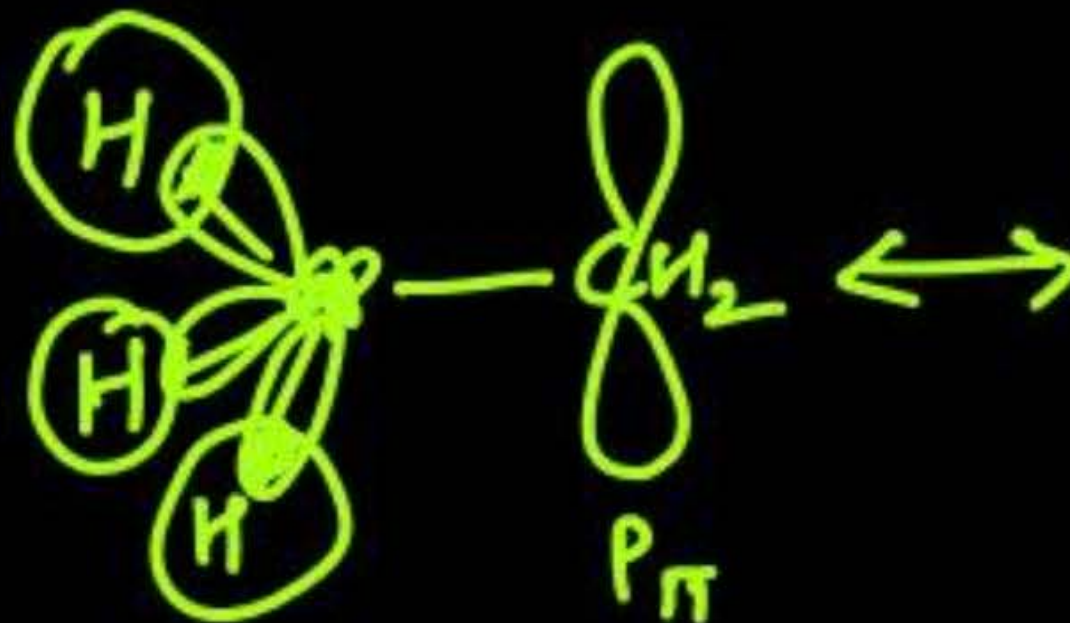
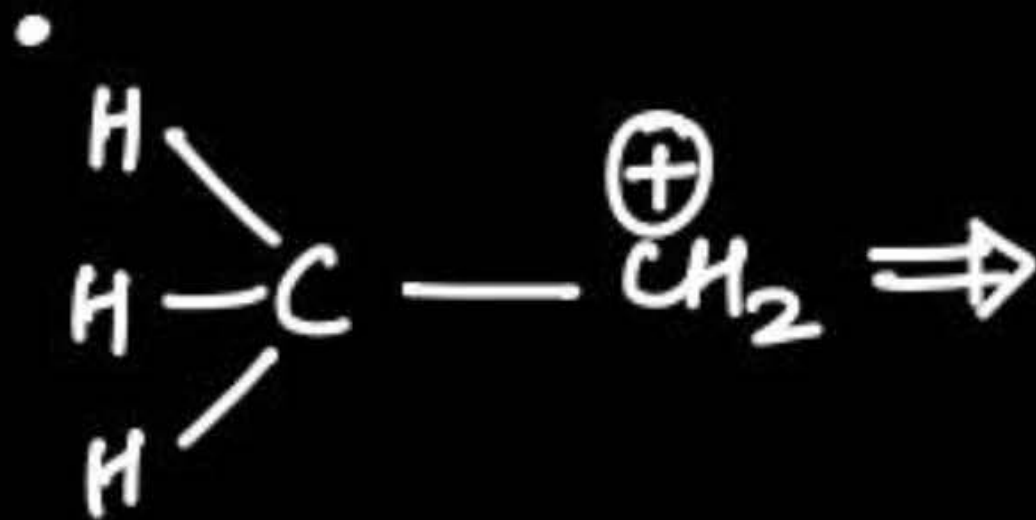
Definition

* When vacant orbital (c^+ , B), alkene, Alkyne are present next to σ bond (C-H) then σe^- of C-H bond delocalised to that vacant orbital. This phenomenon is known as "Hyperconjugation"

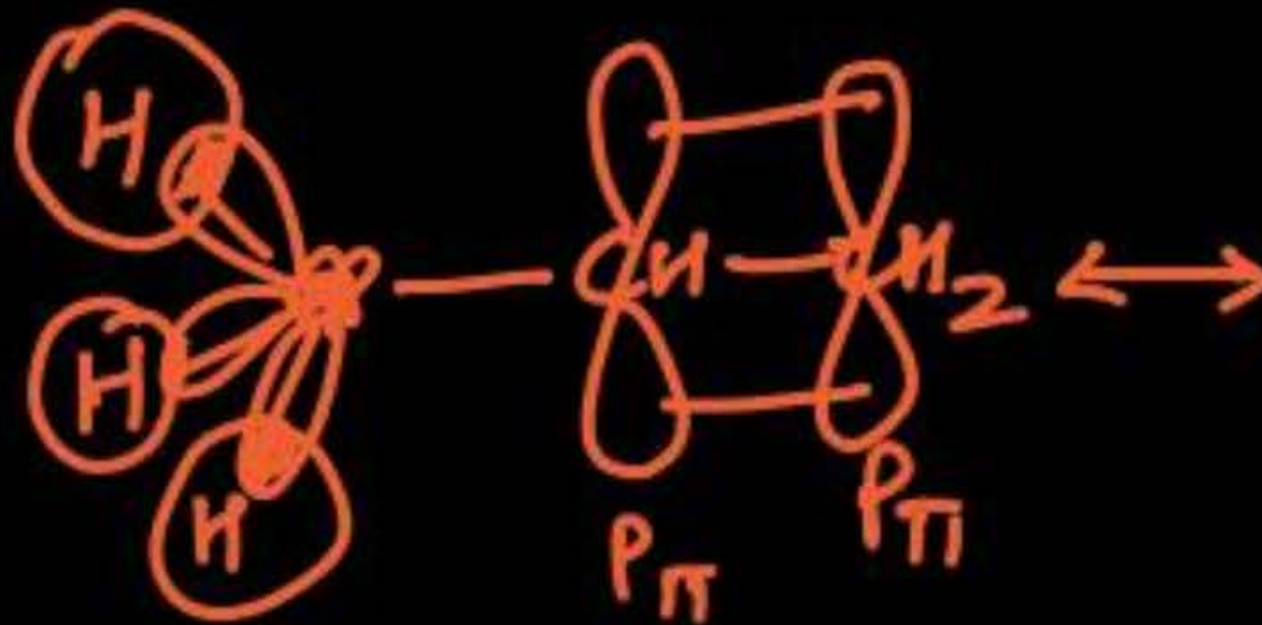
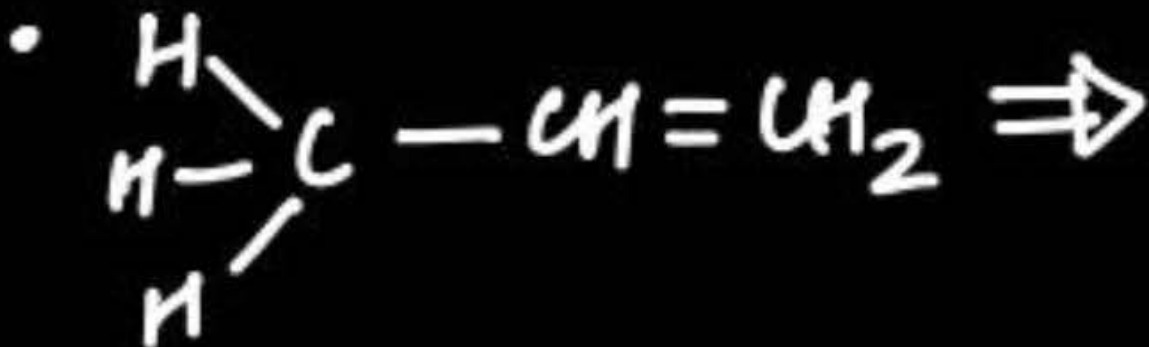
* Delocalisation of σ electrons (C-H) is known as Hyperconjugation.



Presentation



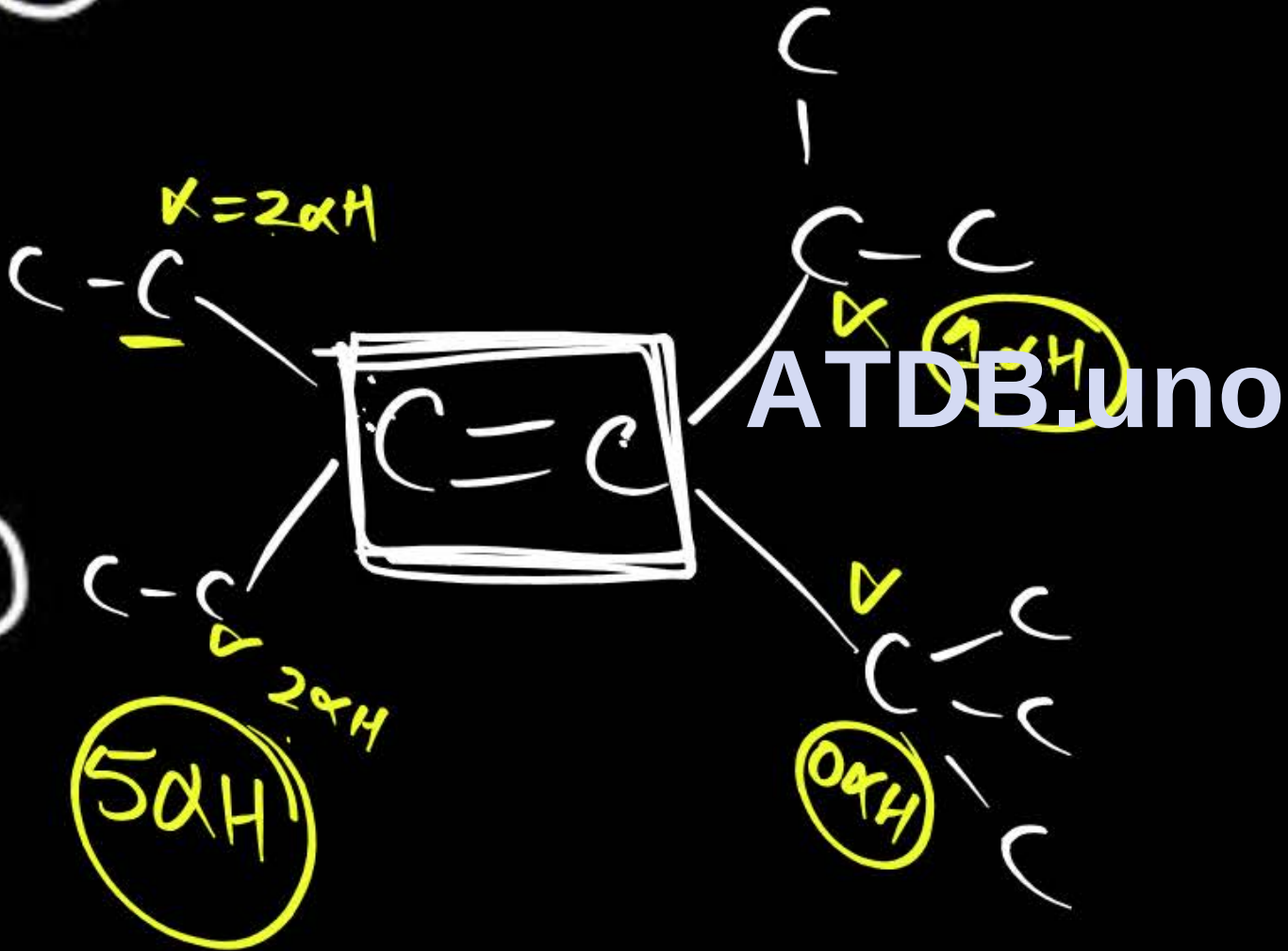
ATDB.uno



Finding of alpha-H

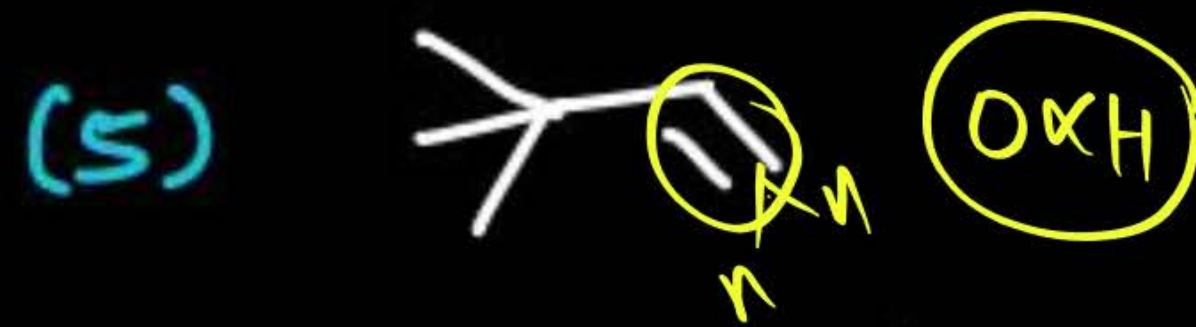
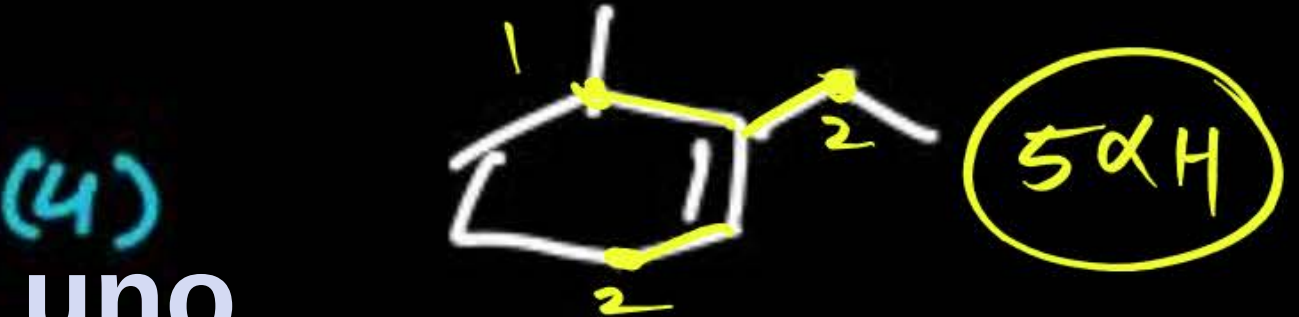
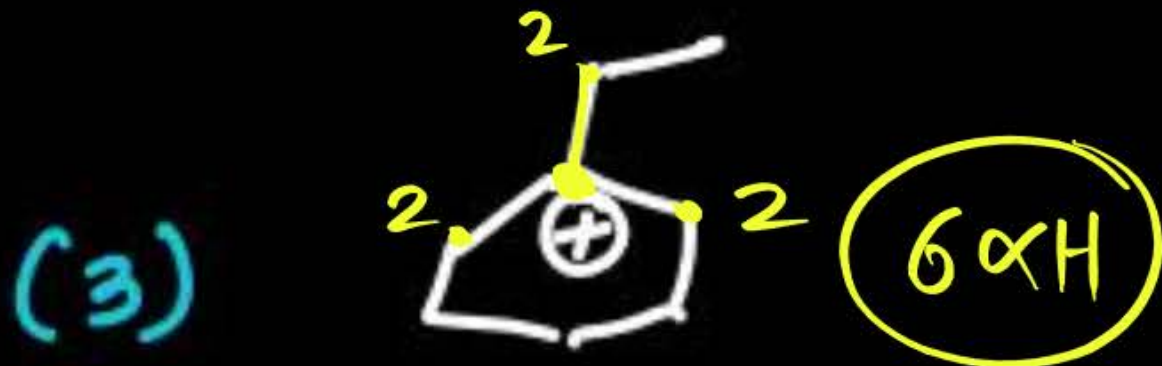
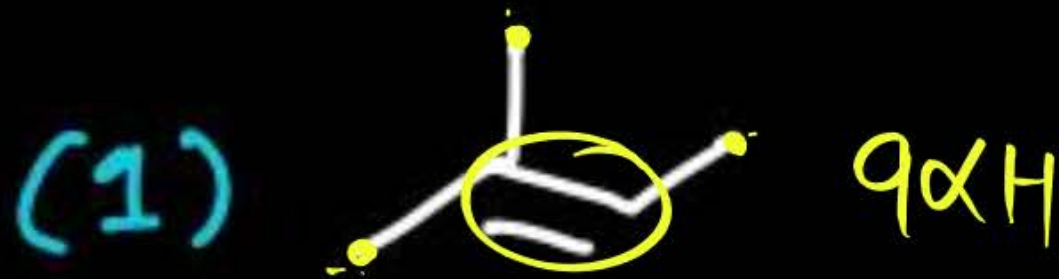


* Cation ✓✓



* Alkene

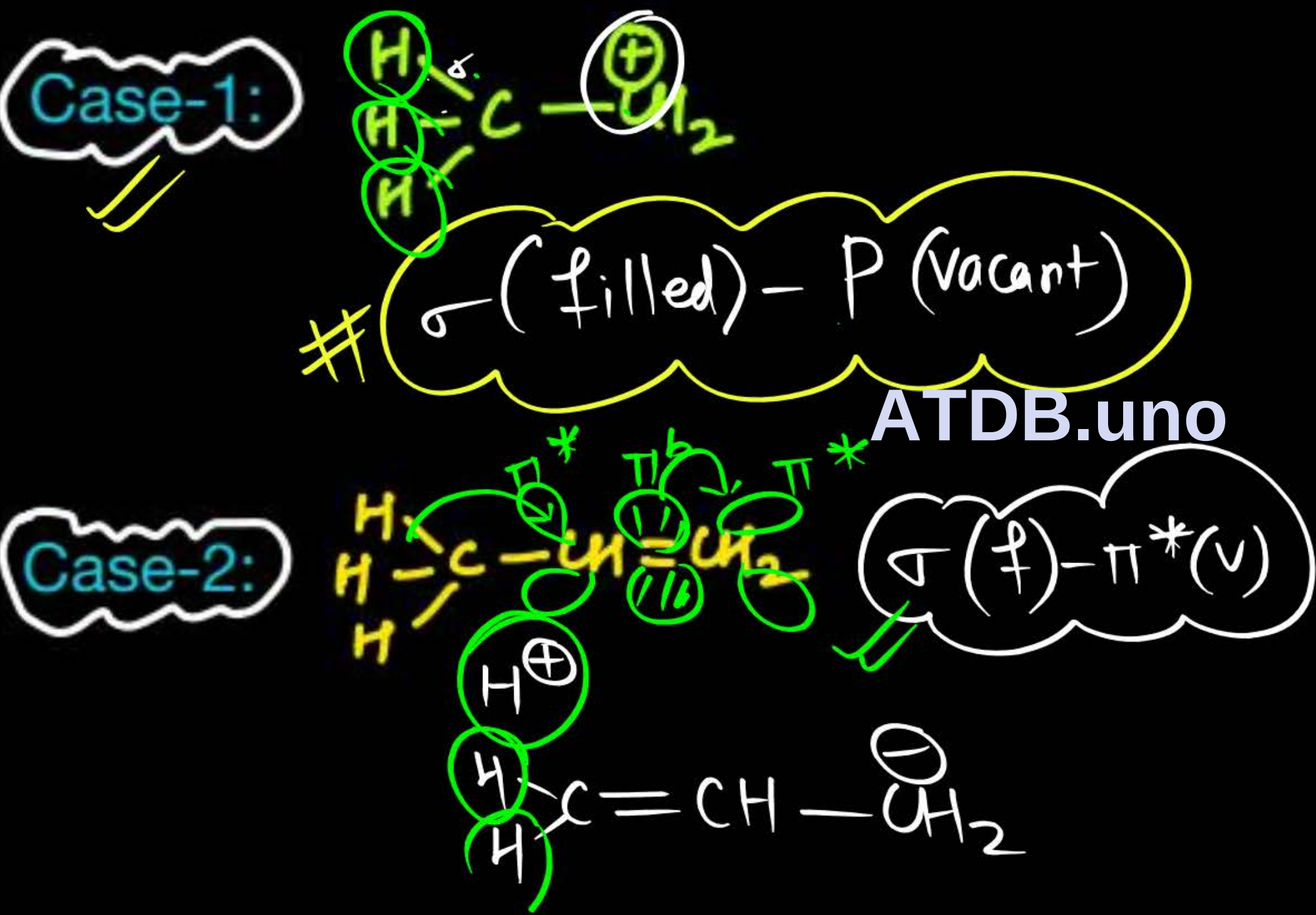
Que Find total alpha H present in following species ?



ATDB.uno



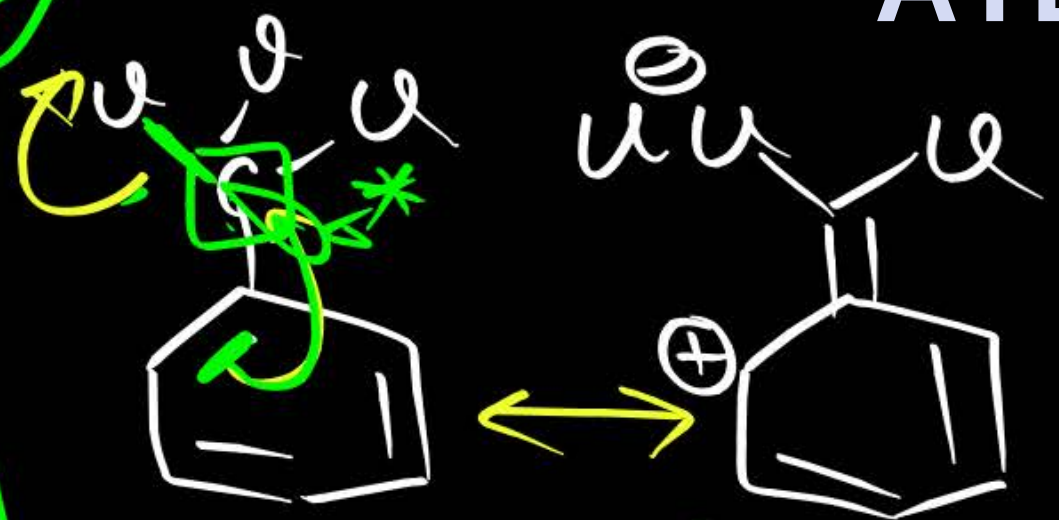
Hyperconjugating Structures



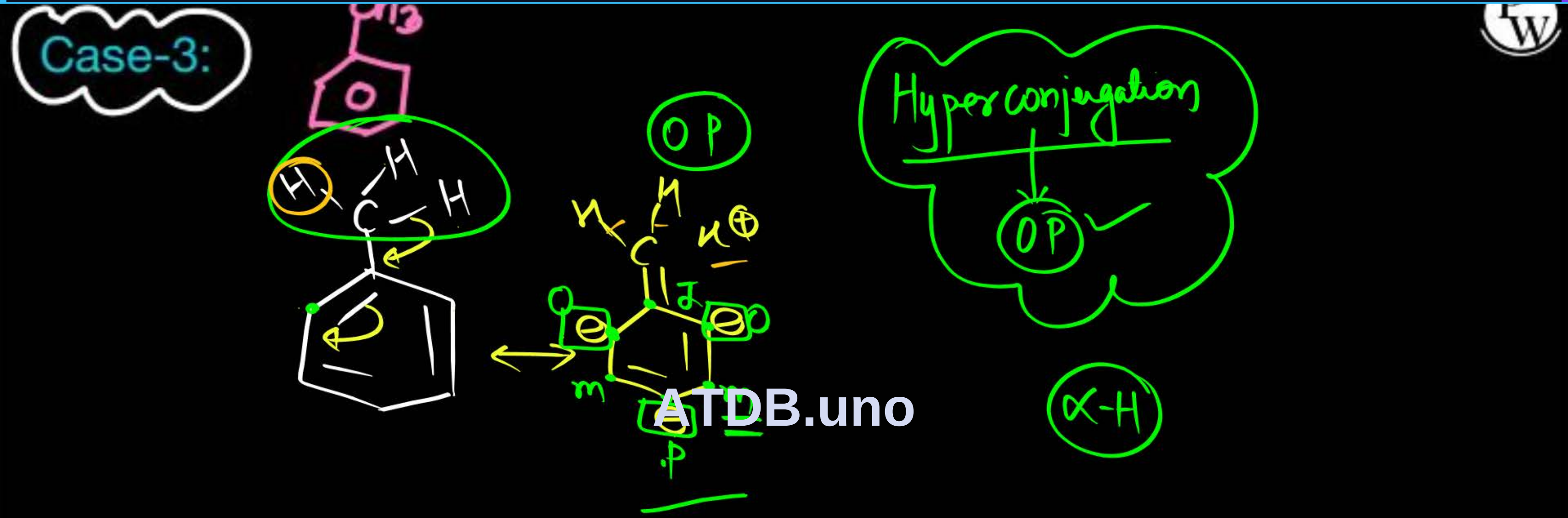


Reverse Hyper (only by CCl_3)

ATDB.uno



π (filled) $\leftrightarrow$ * (vacant)



Note:

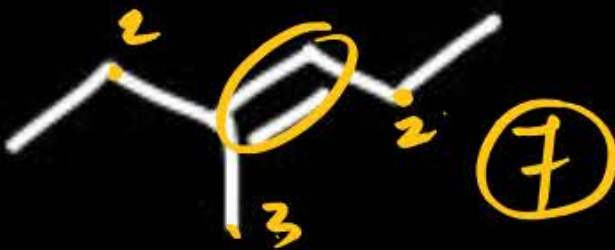
Total H.S. involving C-H bond in benzene = $3 \times \text{no. of } \alpha\text{-H}$

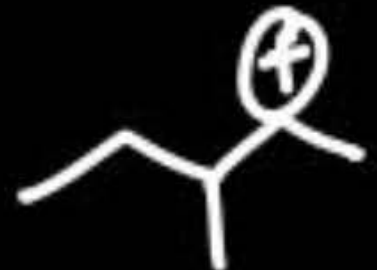
* Total hyperconjugating structures involving C-H bond = no. of $\alpha\text{-H}$

ATDB.uno

* Total hyperconjugating structures = no. of $\alpha\text{-H} + 1$

Q. Find total H.S. Involving C-H bond in following compound ?

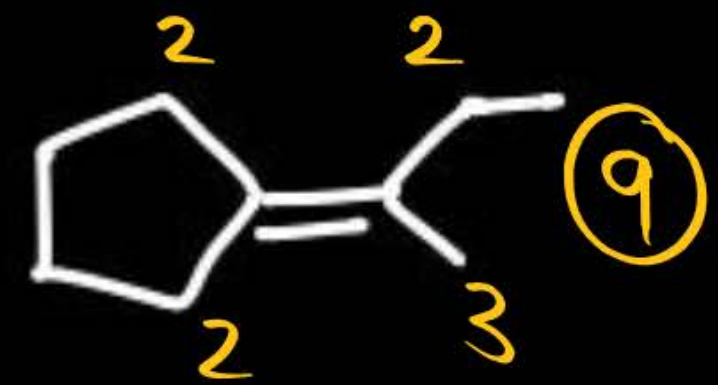
(1) 

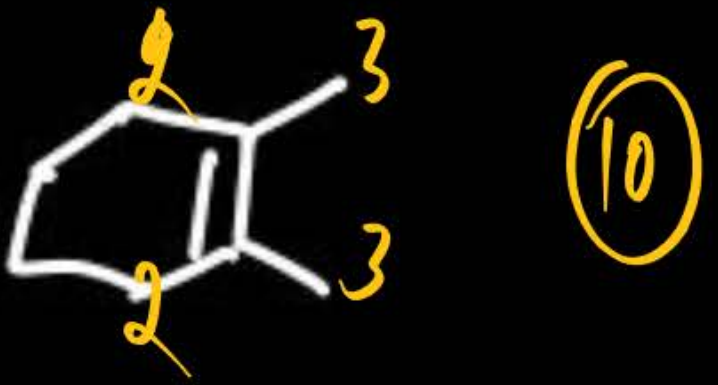
(2) 

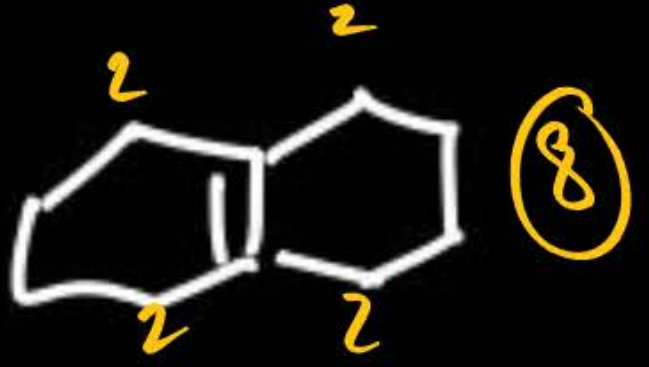
(3) 

(4) 

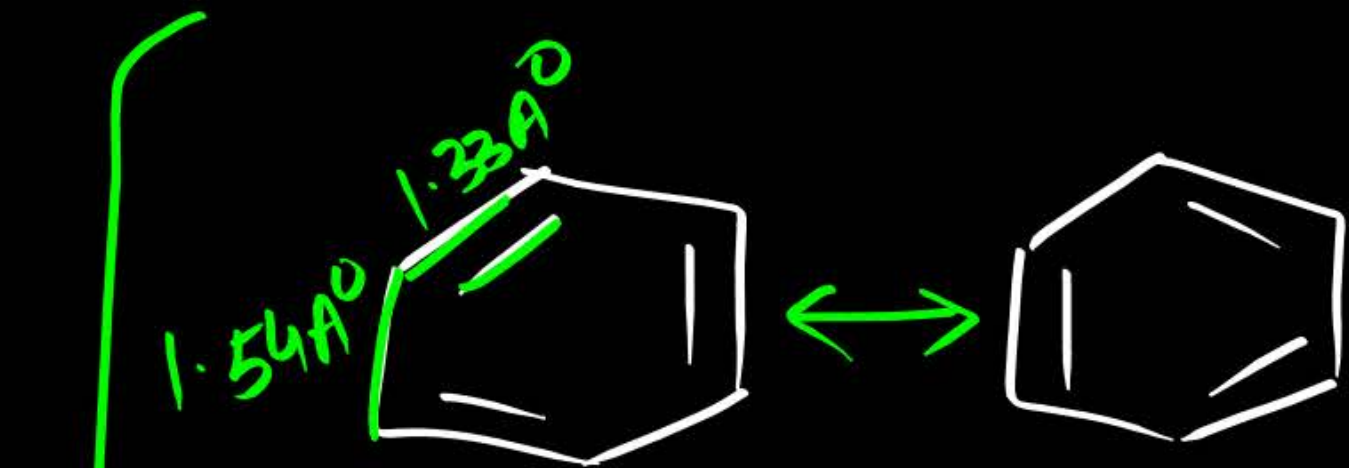
(5) 

(6) 

(7) 

(8) 

ATDB.uno



* Hypothetical

* Resonating
Str. or



Real

* Hybrid

ATDB.uno

Definitions



- When properties of a compound can't be explained by a single structure then various structures are drawn.
- These structures are known as resonating structures or canonical structures or contributing structures while real compound is known as resonating hybrid.

ATDB.uno

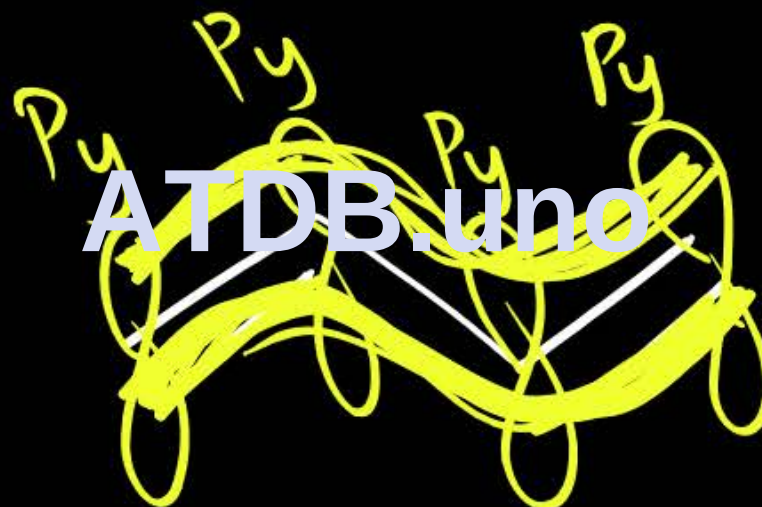
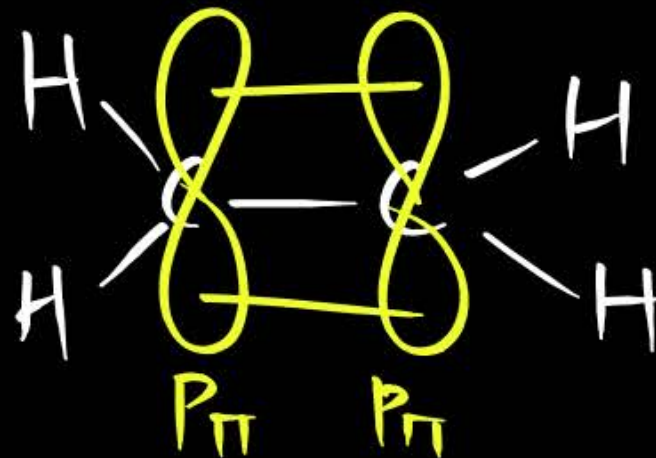
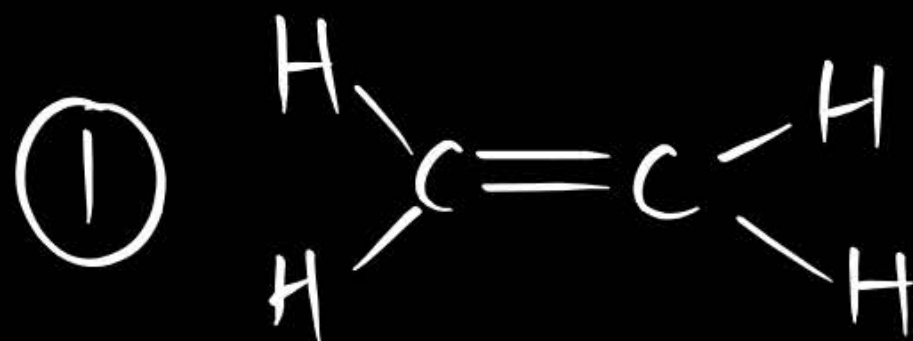
● Ex:



Properties Of Resonance

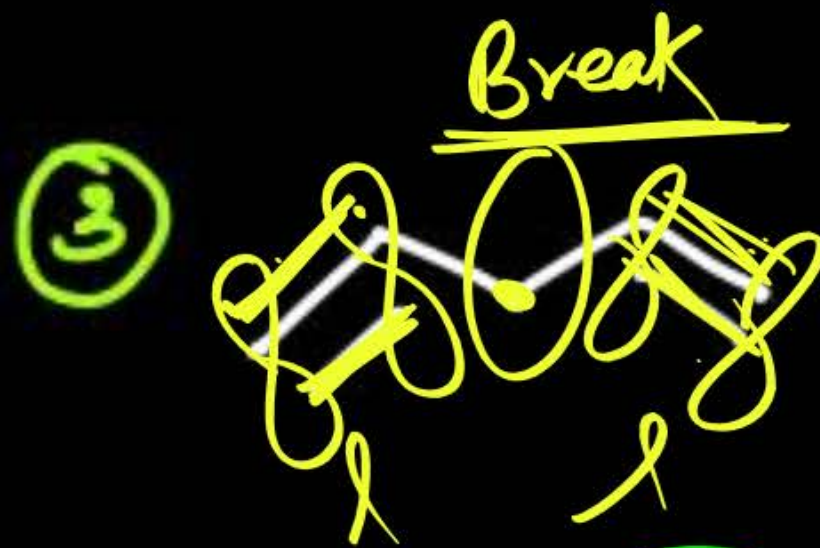
- It is delocalisation of pi electrons.
- It is a permanent effect and strongest effect.
- It is intramolecular phenomenon.
- Generally most stable RS contribute most.
- It is a distance independent effect.
- Resonating structures are hypothetical and hybrid is real.

ATDB.uno



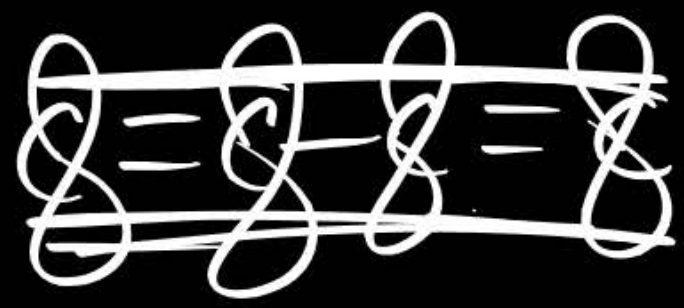
#

$a = \text{no. of } p_{\pi} = 4$
 $b = \text{no. of } p_{\pi} \text{ in Reso} = 4$
 $c = \text{no. of } \pi e^{-} \text{ in Reso} = 4$



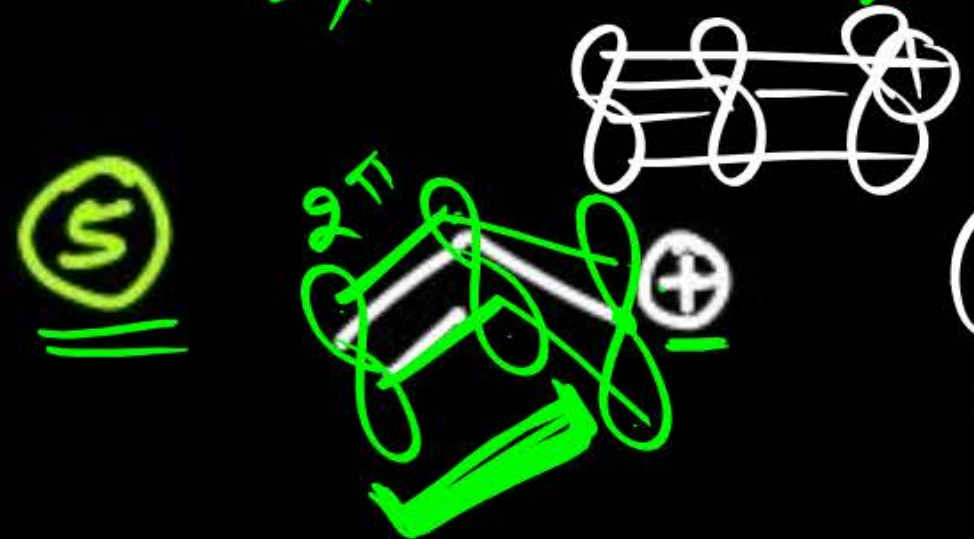
Reso X

$a = 4$
 $b = 0$
 $c = 0$



$a = 6$
 $b = 4$
 $c = 4$

ATDB.uno



$a = 3$
 $b = 3$
 $c = 2$

no. of π e⁻ =

6



$a=6$
 $b=6$
 $c=6 \rightarrow A$

7



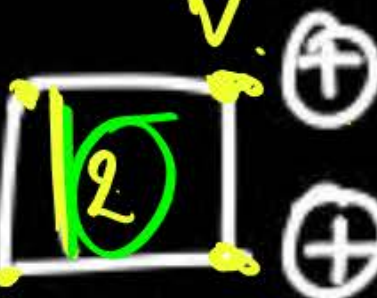
$a=4$
 $b=4$
 $c=4 \rightarrow AA$

8



$a=4$
 $b=4$
 $c=4 \Rightarrow NA$

9



$a=4$
 $b=4$
 $c=2$

A

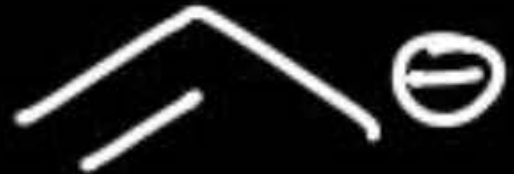
10



$a=7$
 $b=7$
 $c=6$

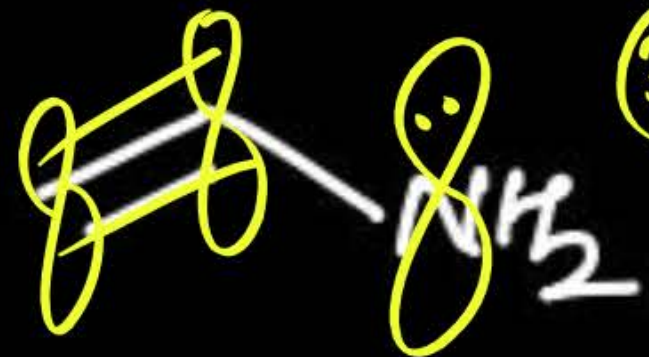
A

11



$(3,3,4)$

12



$(3,3,4)$

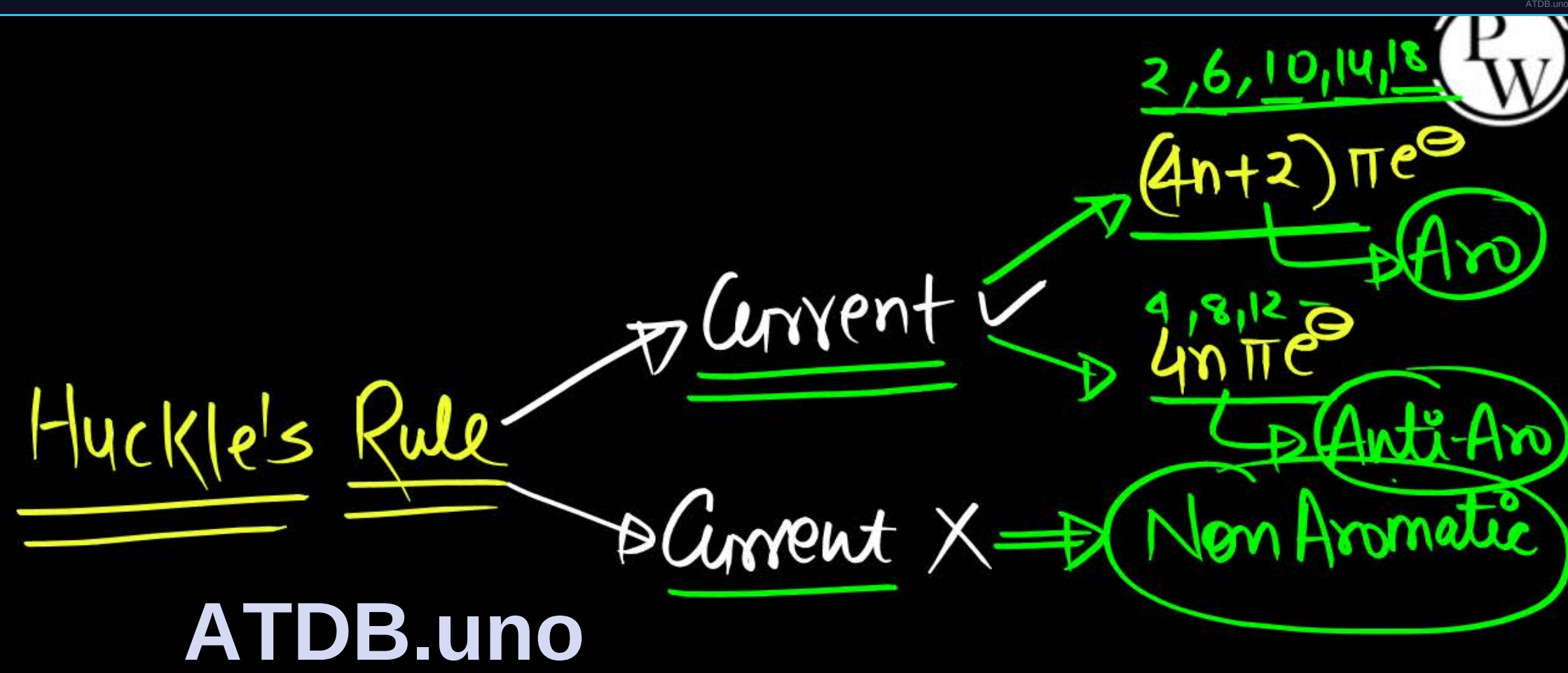
13



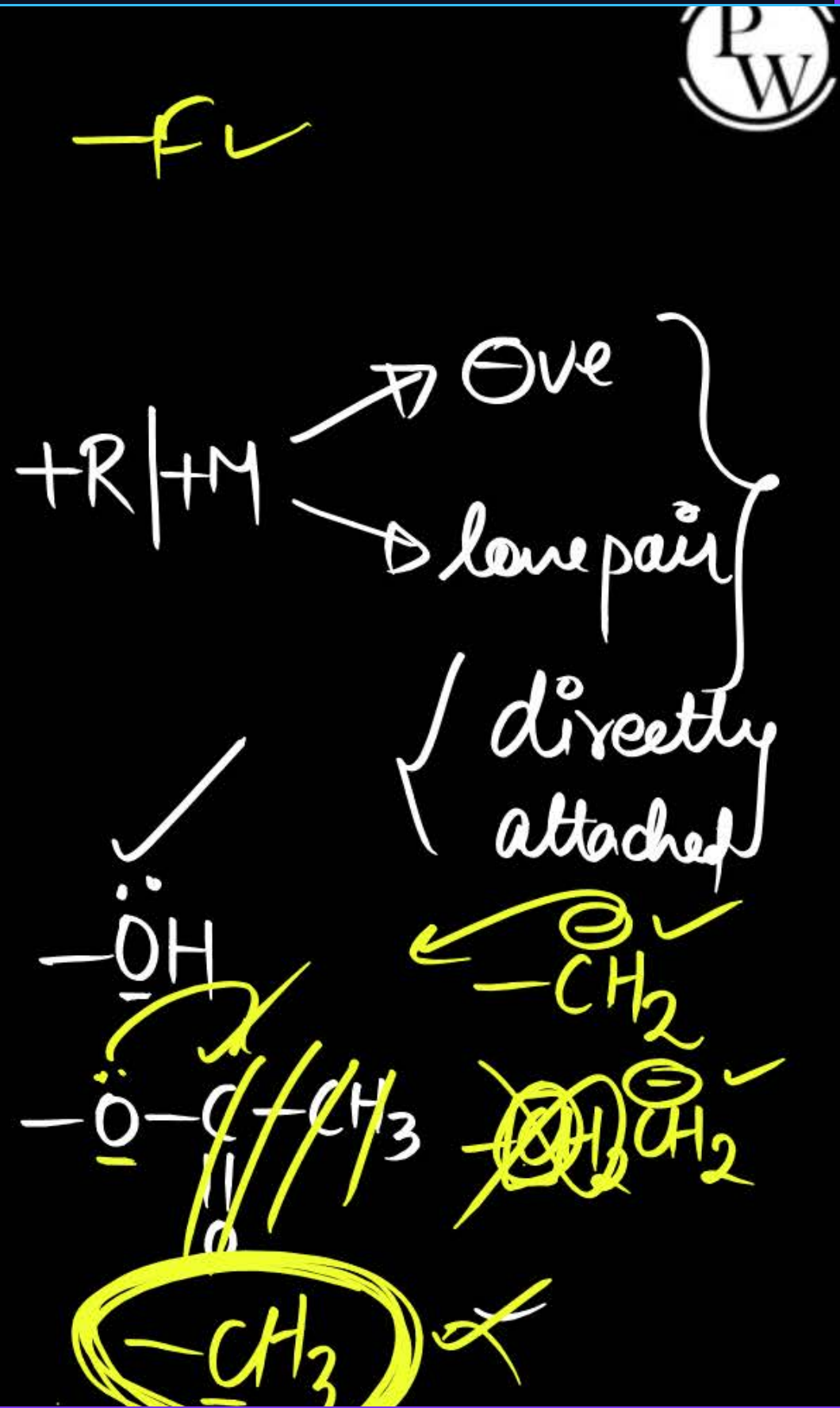
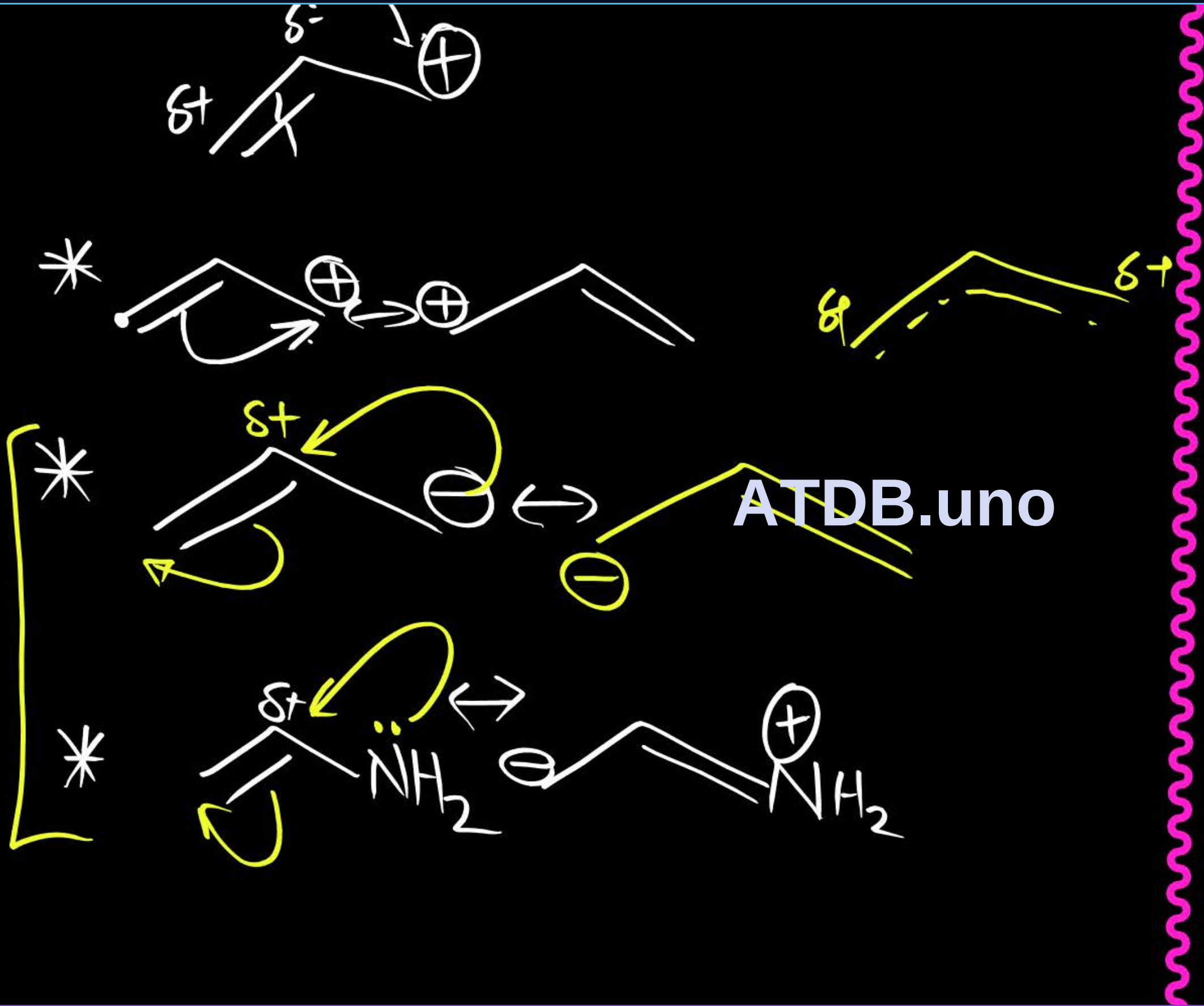
$(3,3,2)$

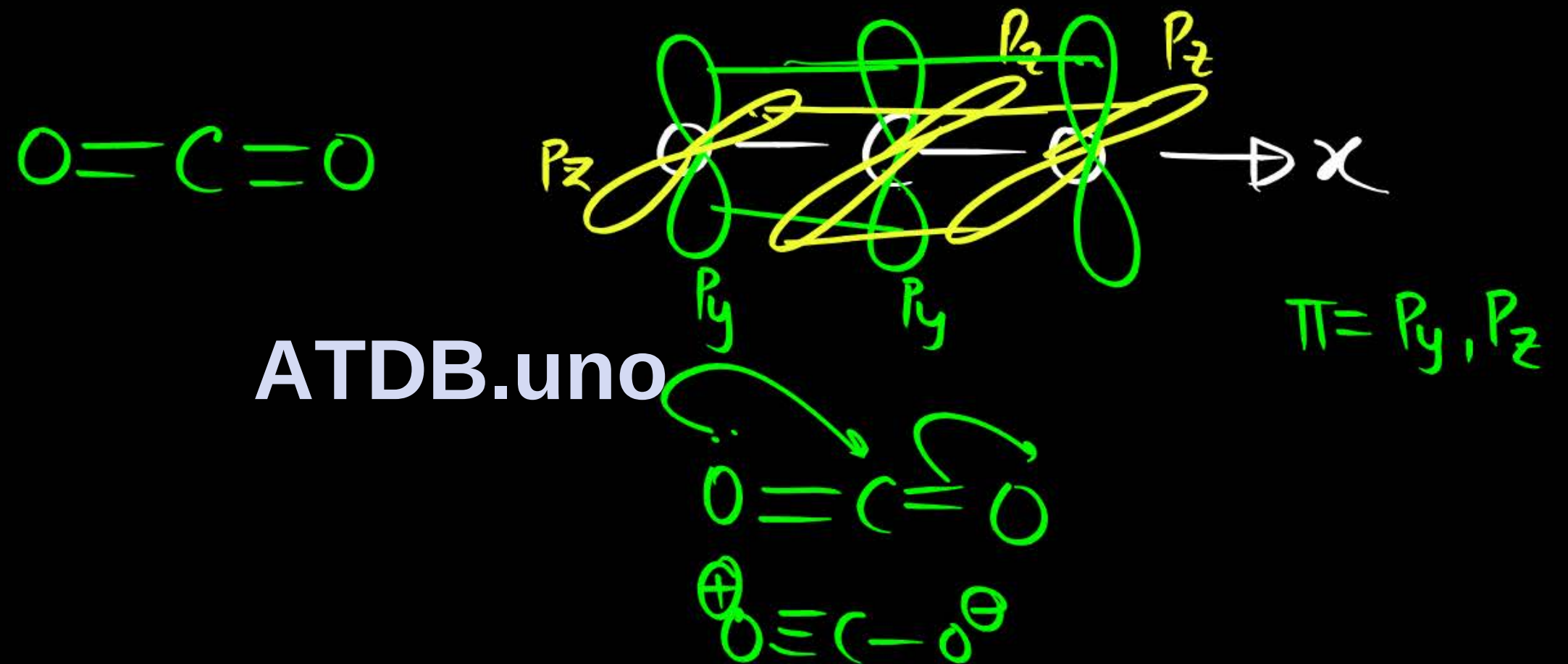
14





ATDB.uno





ATDB.uno



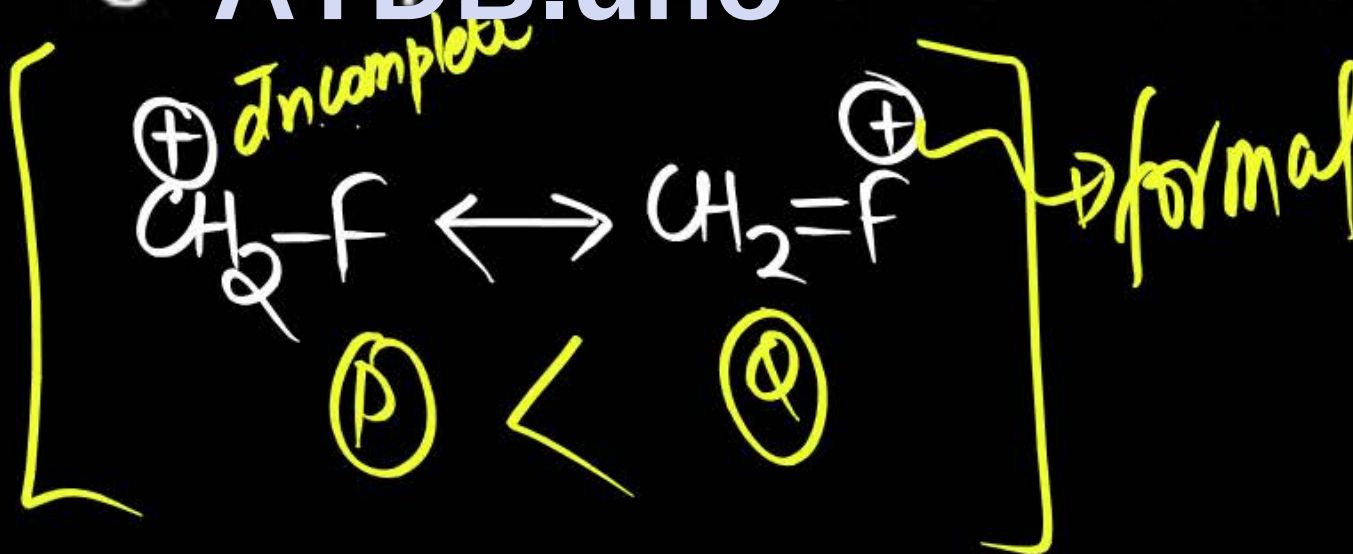
- * Octet
- * Neutral
- * Over on move EN
- * $\ominus | \oplus$
- * fries
- * defect



Stability Of RS

Rule-1 Octet

RS having complete octet is more stable





Rule-2 Neutral

Generally Neutral RS is more stable than ionic

ATDB.uno



Rule-3 Electronegative

-ve on More EN atom is more stable and +ve on less EN atom

ATDB.uno



Rule-4 Opposite Charges

RS having closer opposite charges are more stable as they attract each other

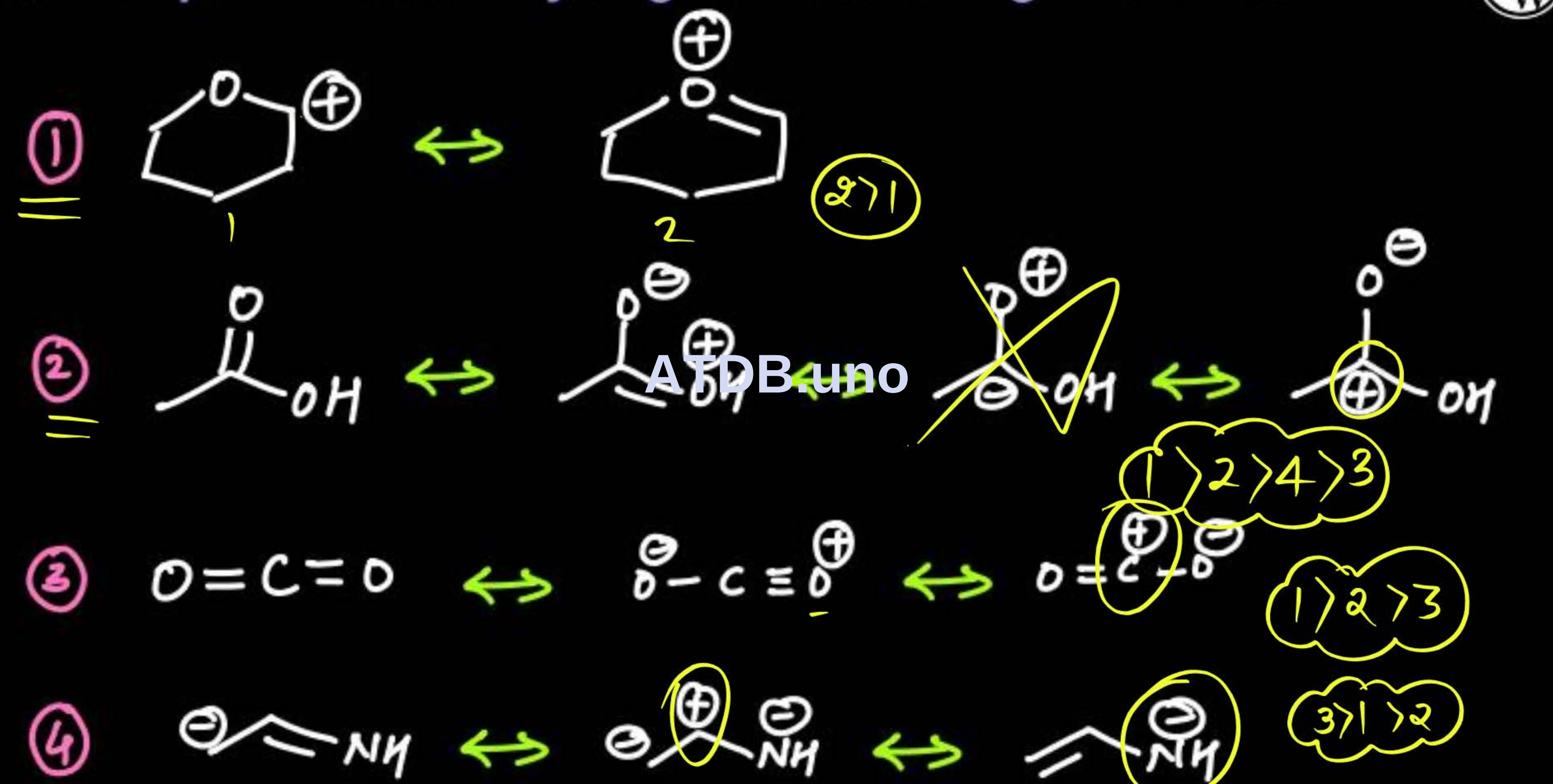
ATDB.uno

Rule-5 Fries Rule

RS having more benzene is more stable

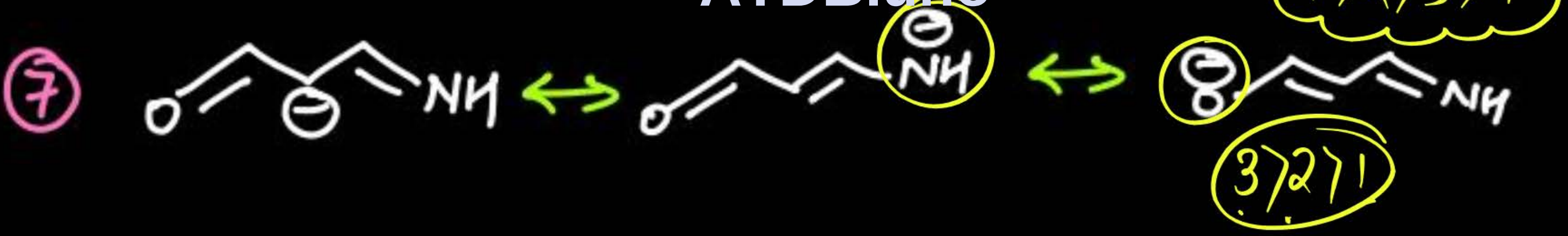
ATDB.uno

Q. Compare the stability of given resonating structures ?

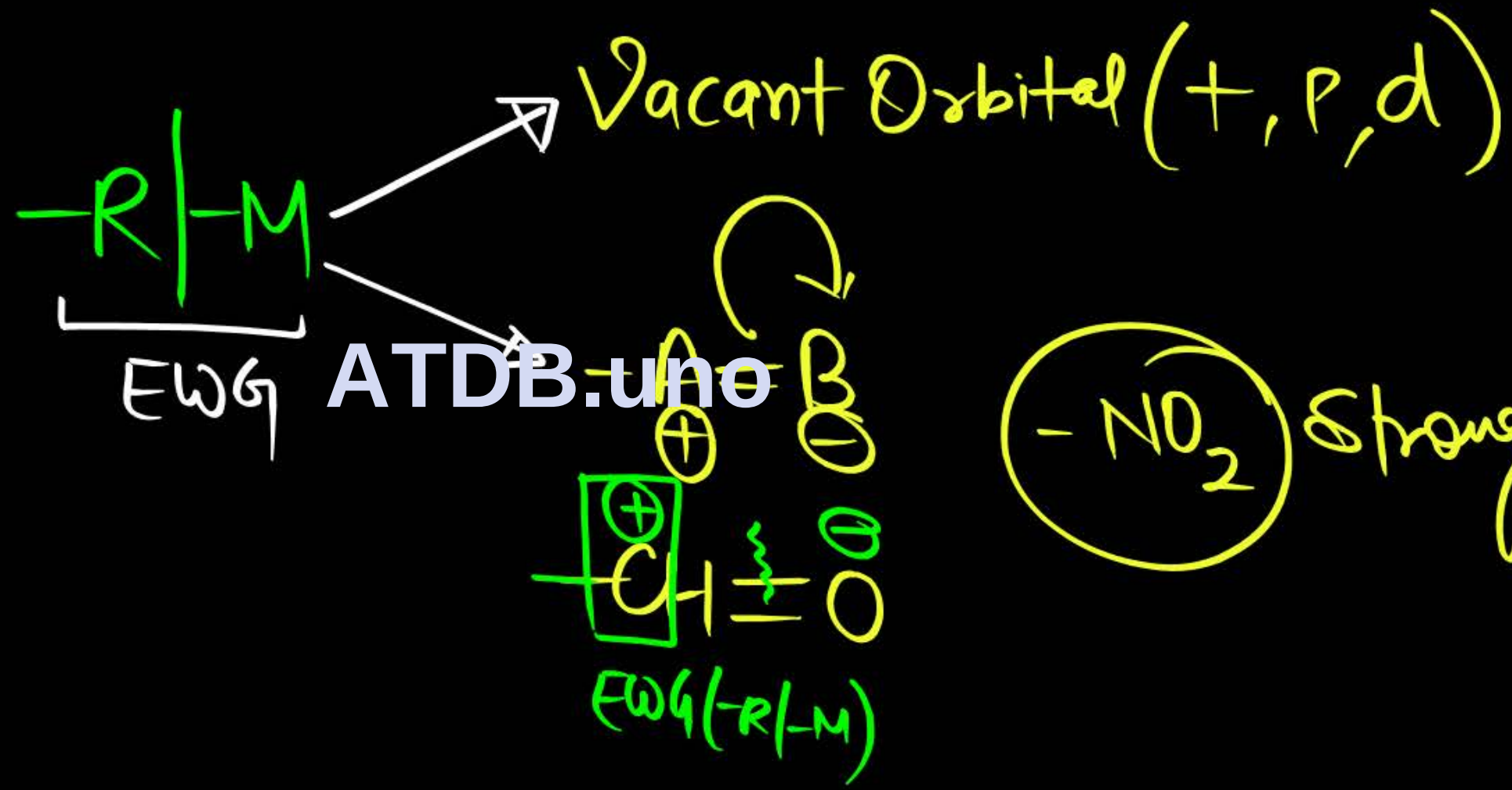




ATDB.uno



Resonance or Mesomeric Effect

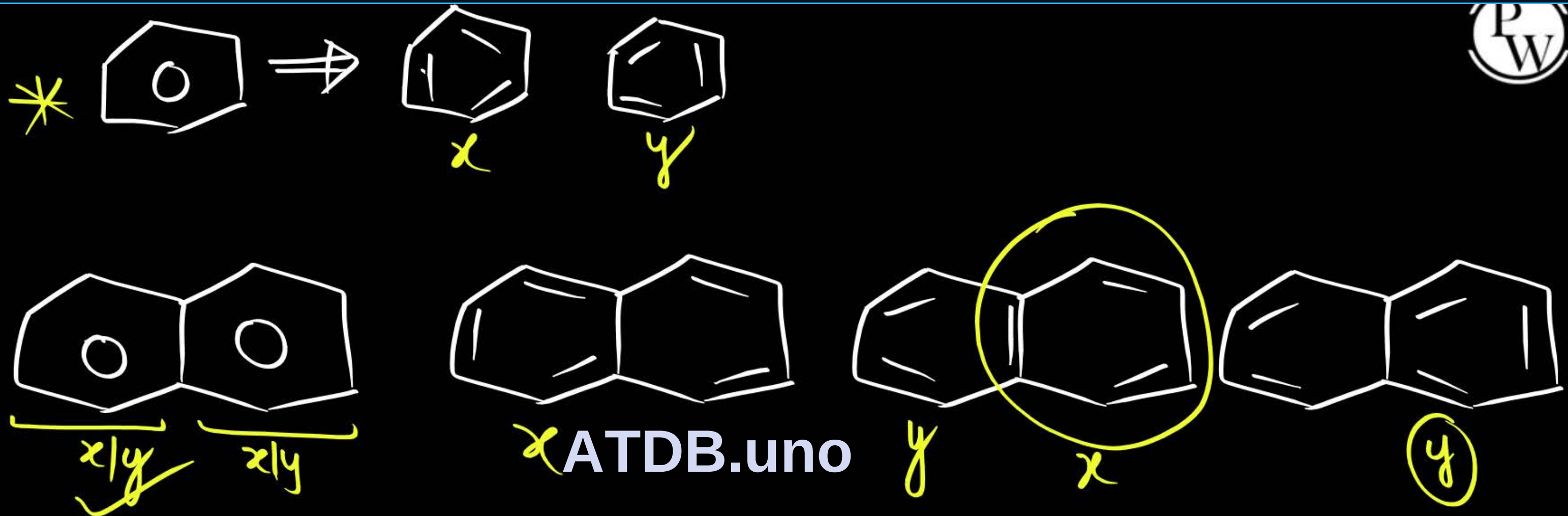


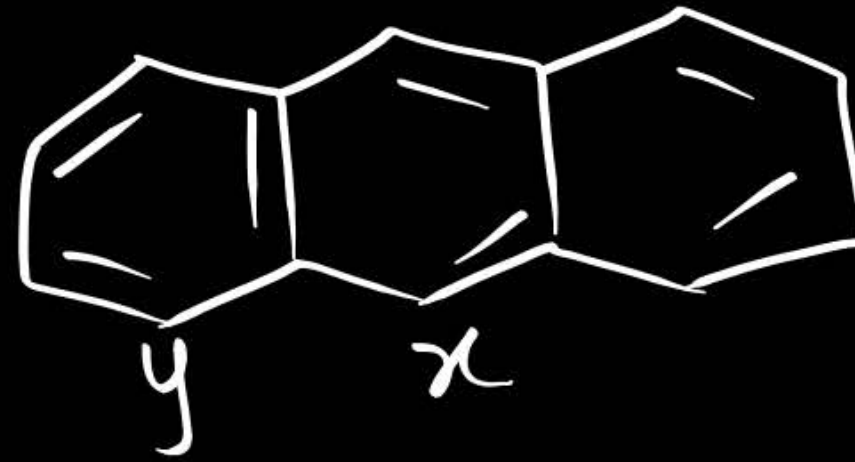
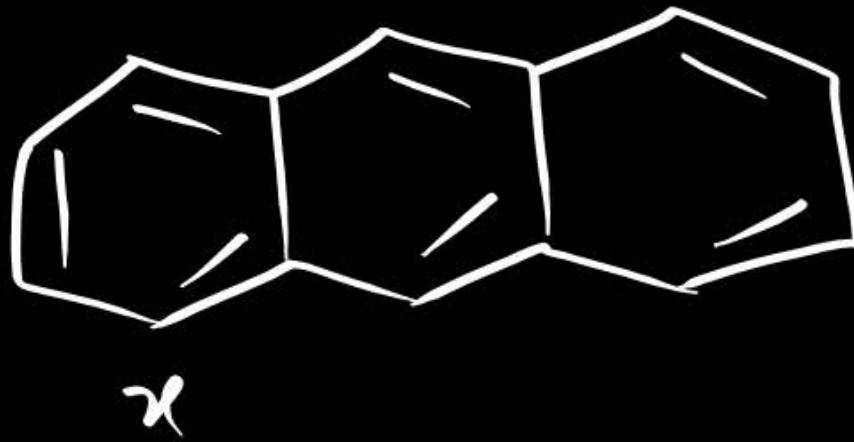
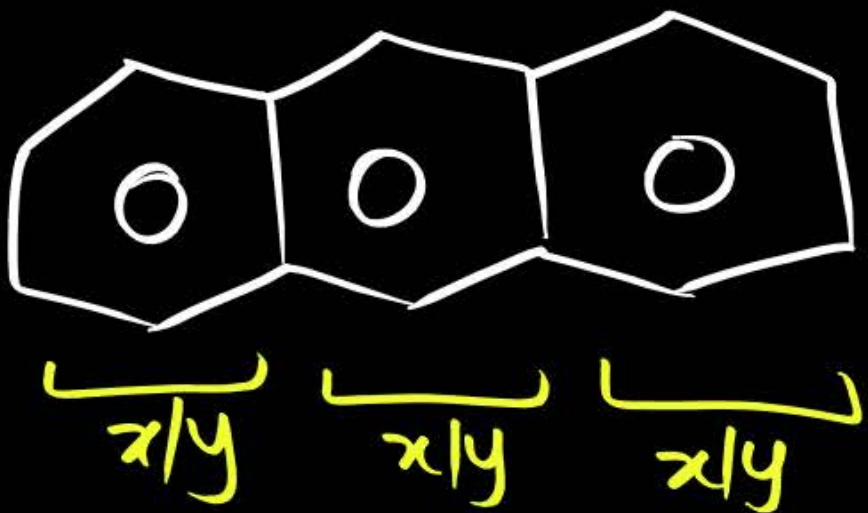
NO_2 Strongest



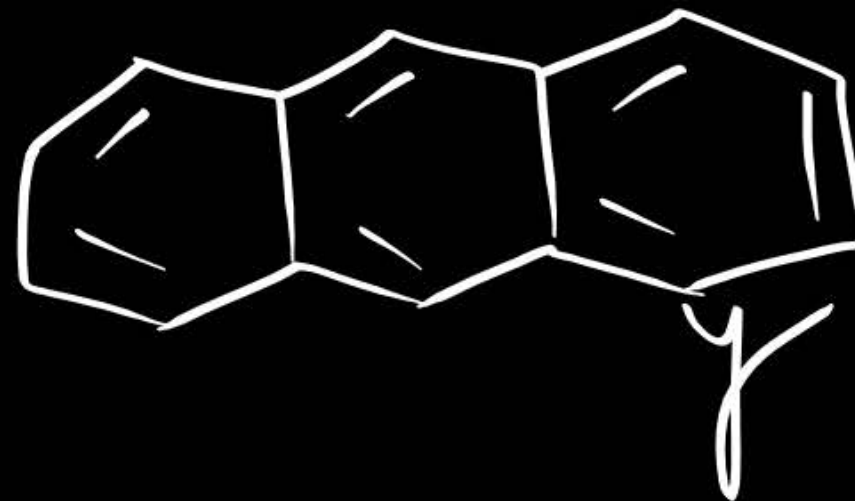
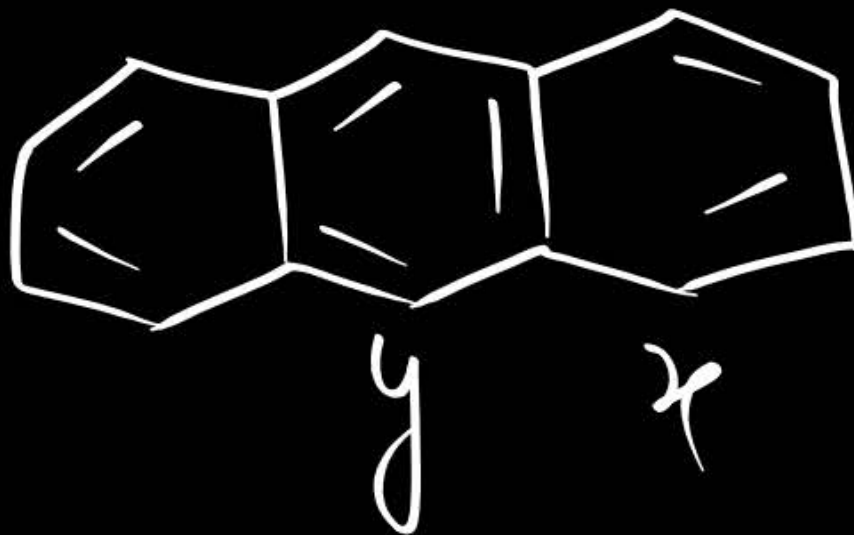
* all Robinhood net shows (-2) with Benzene

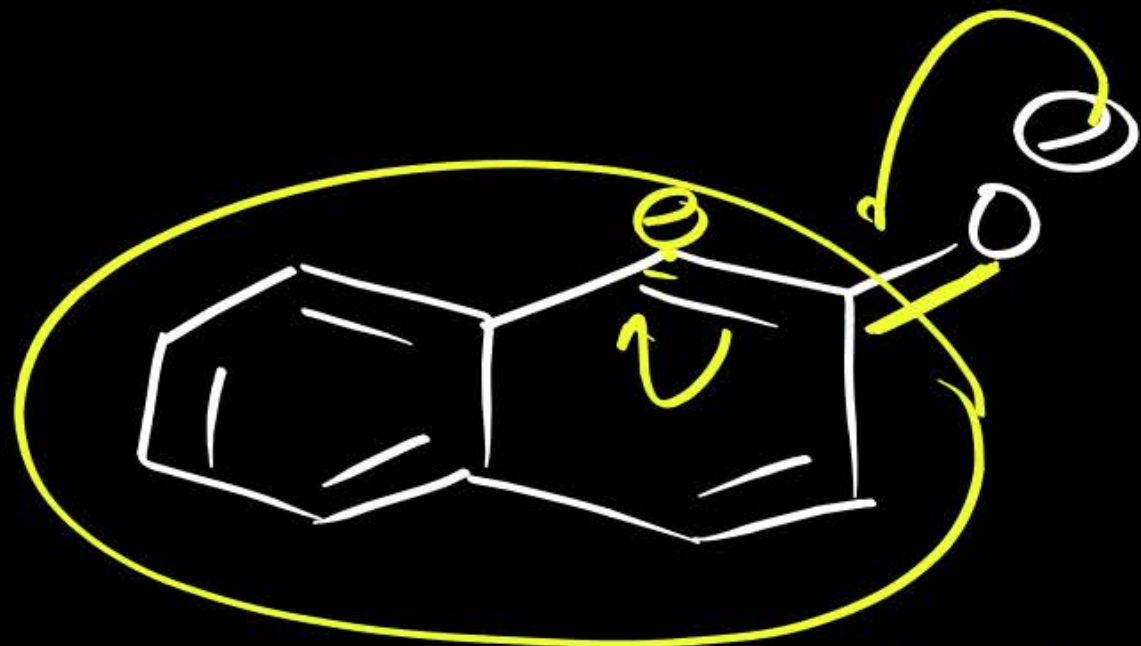
ATDB.uno



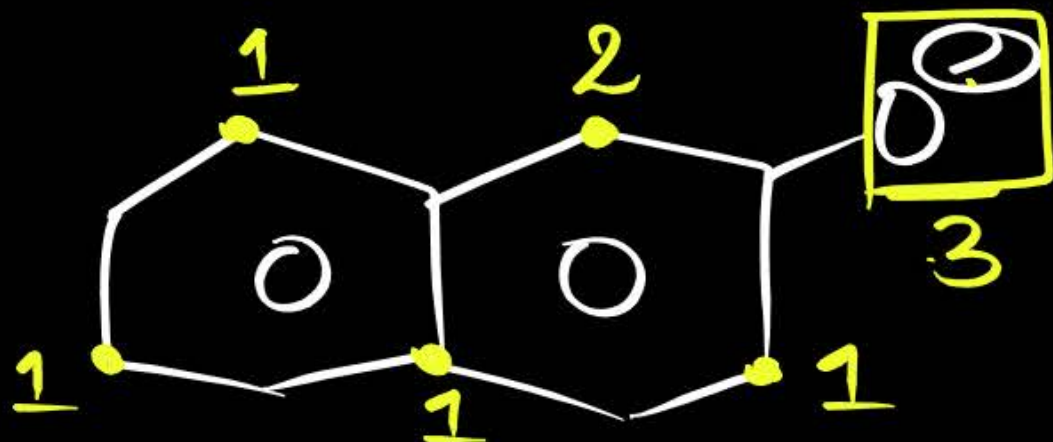


ATDB.uno

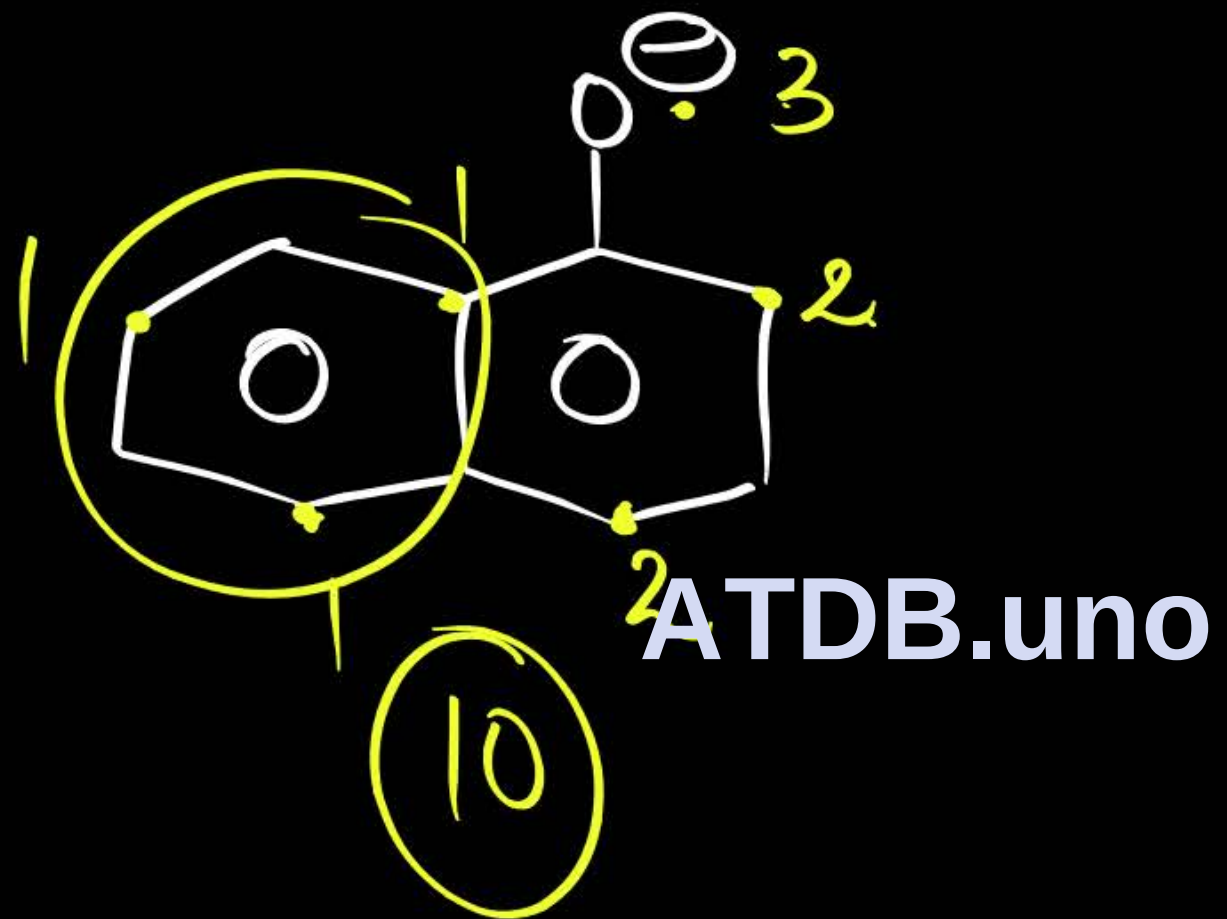




ATDB.uno
* naphthalene



9 ✓



21



(A)

22



(A)

23



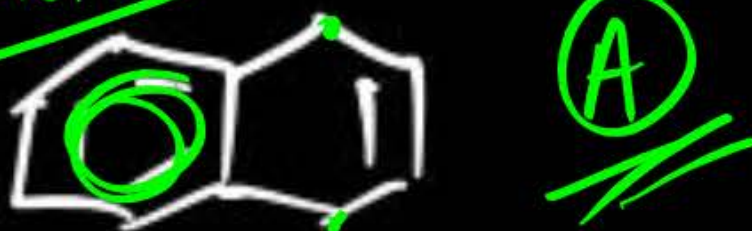
(AA)

24



(A)

25



(A)

26



(A)

27



(A)

28



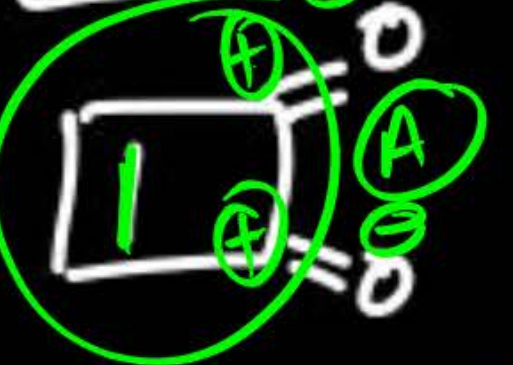
(A)

29



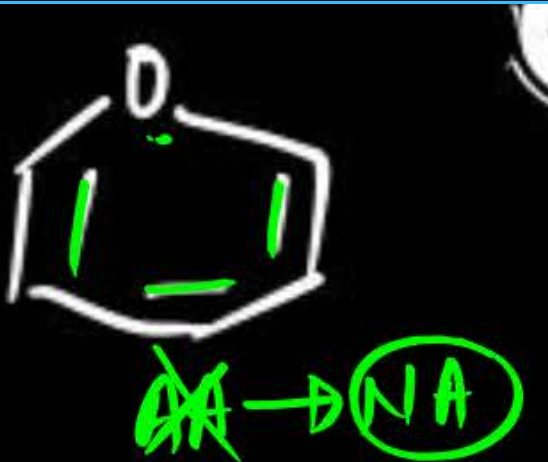
(AA)

30



(A)

31



(NA)

32



(NA)

33



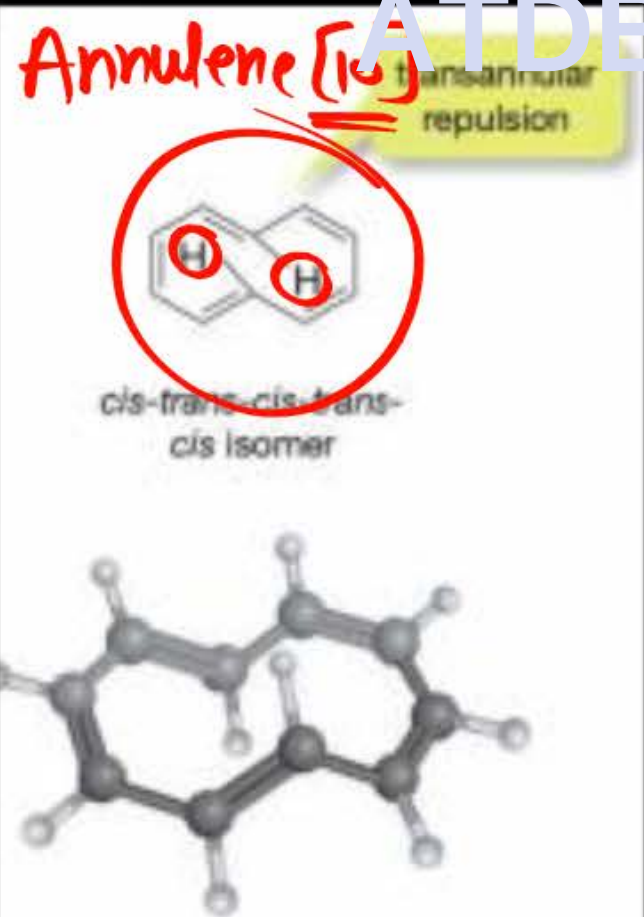
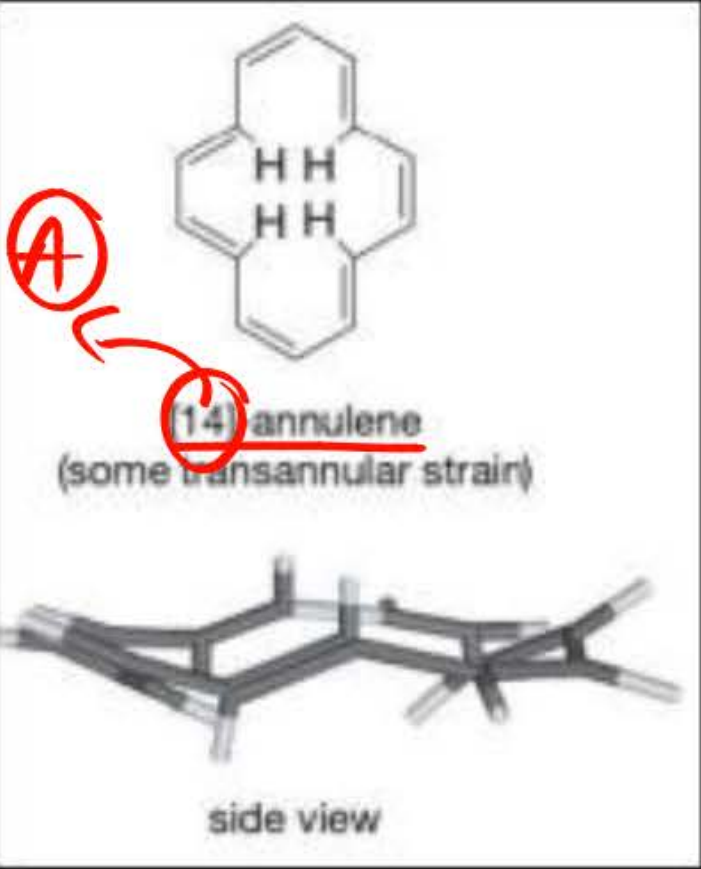
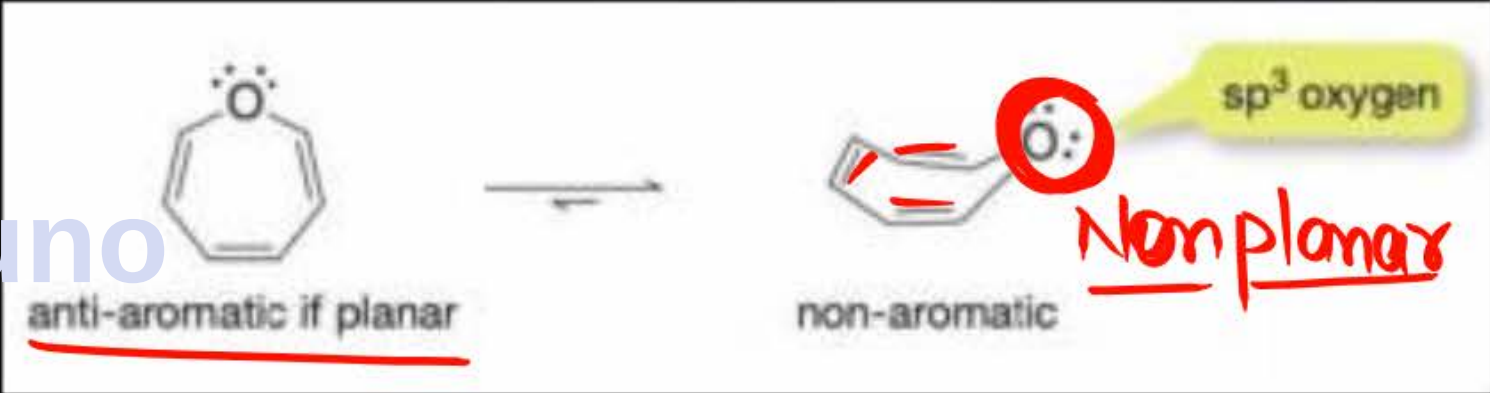
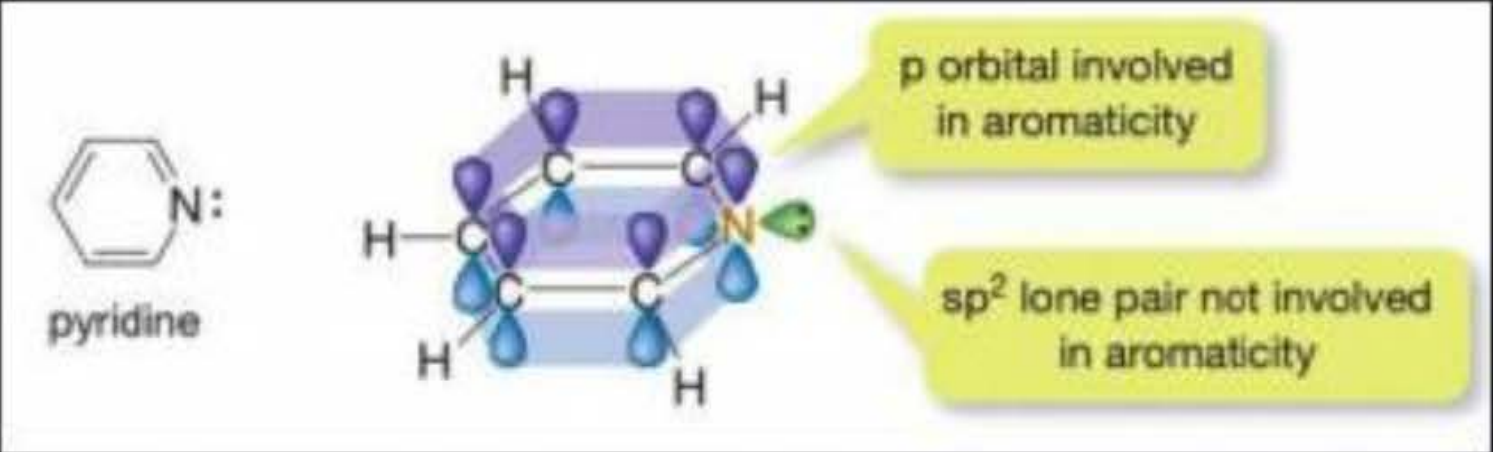
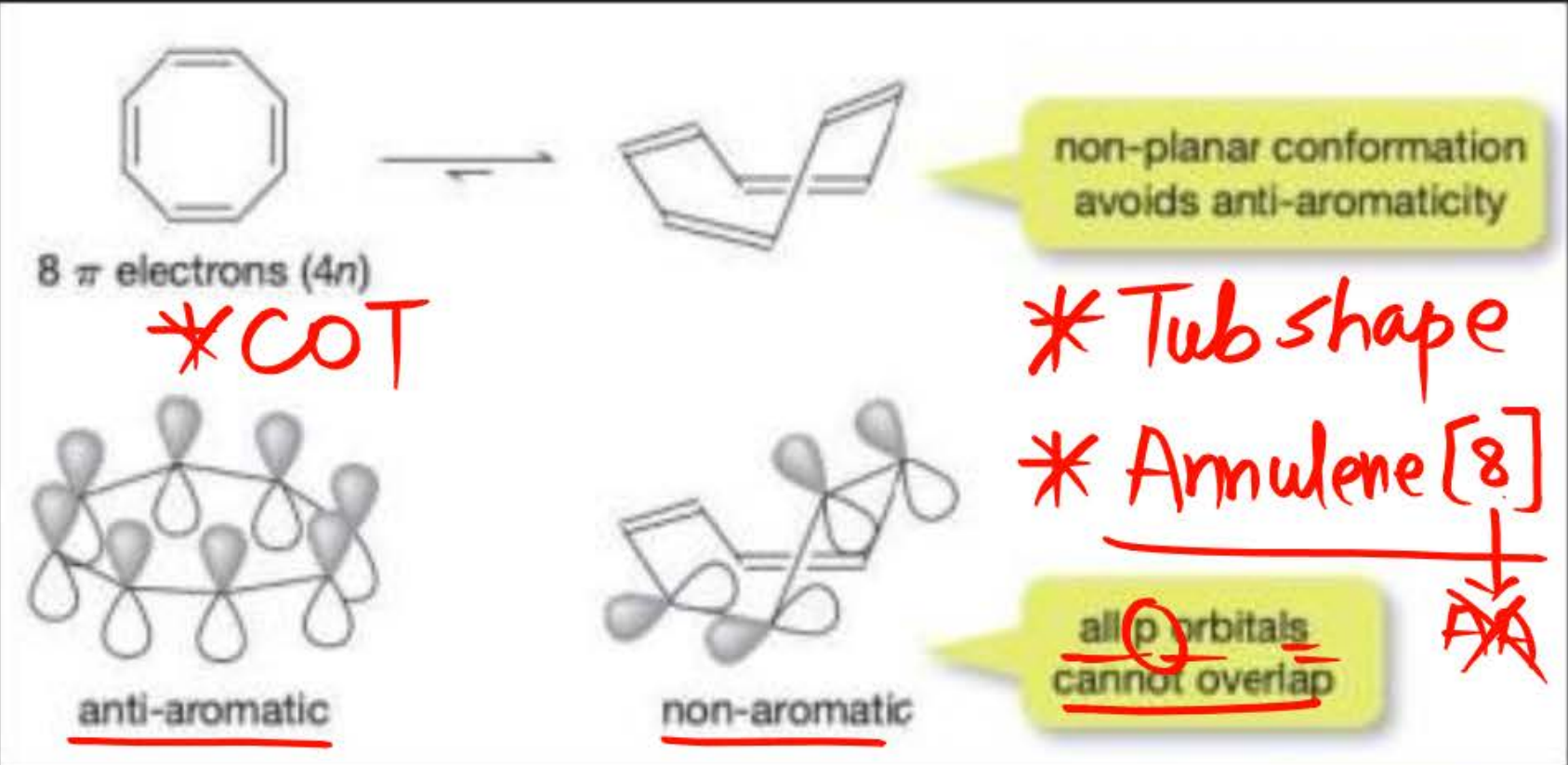
(NA)

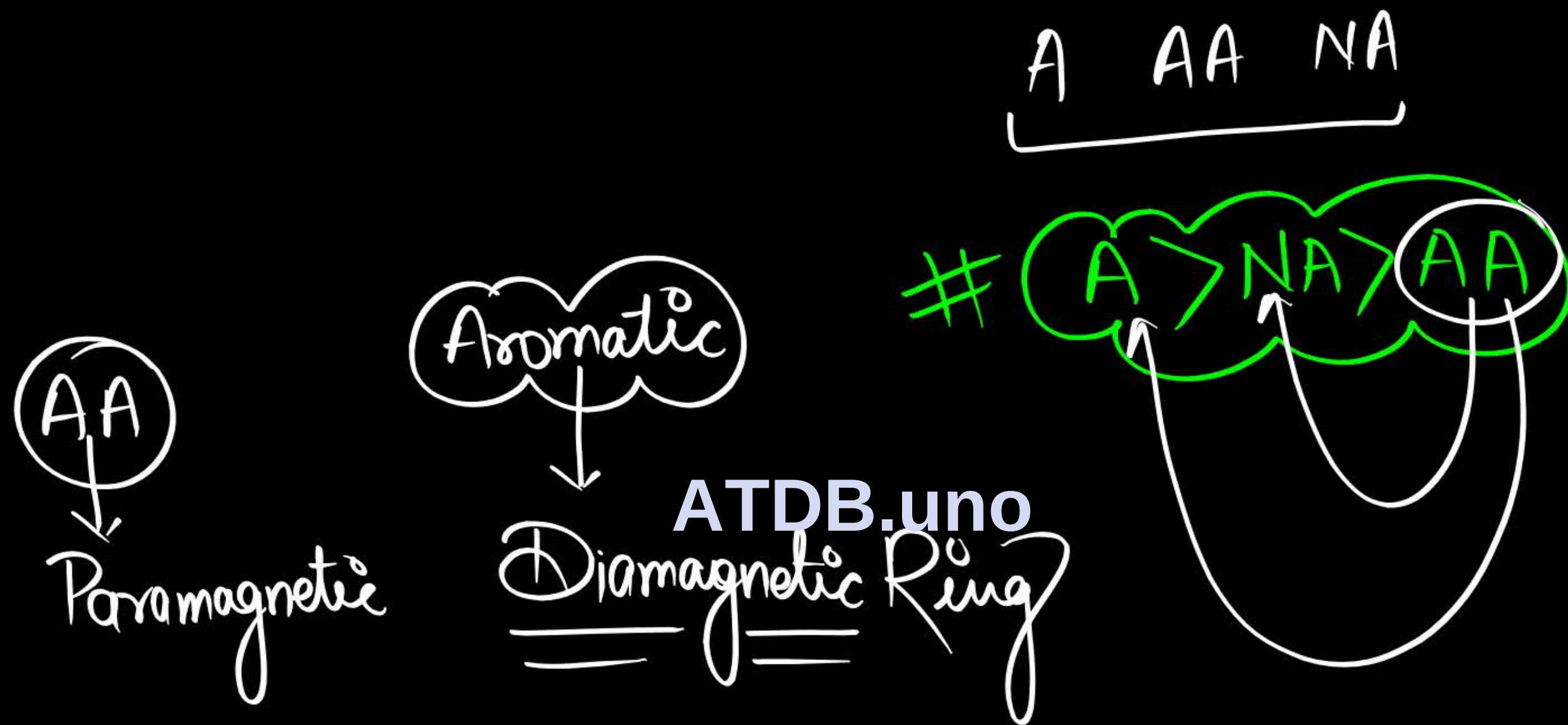
Downloaded from ATDB.uno/StudyPrayas | Unauthorised redistribution is strictly prohibited.

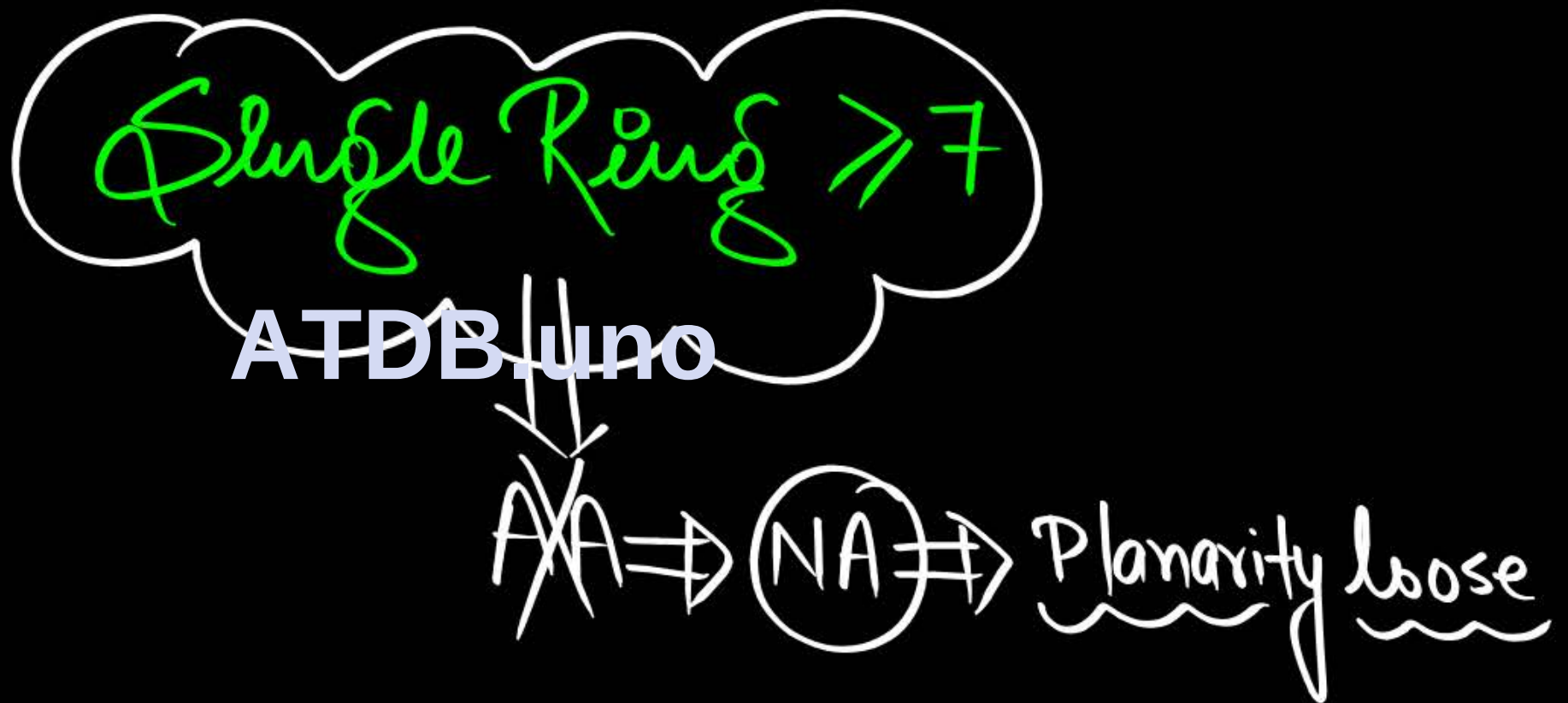
ATDB.uno

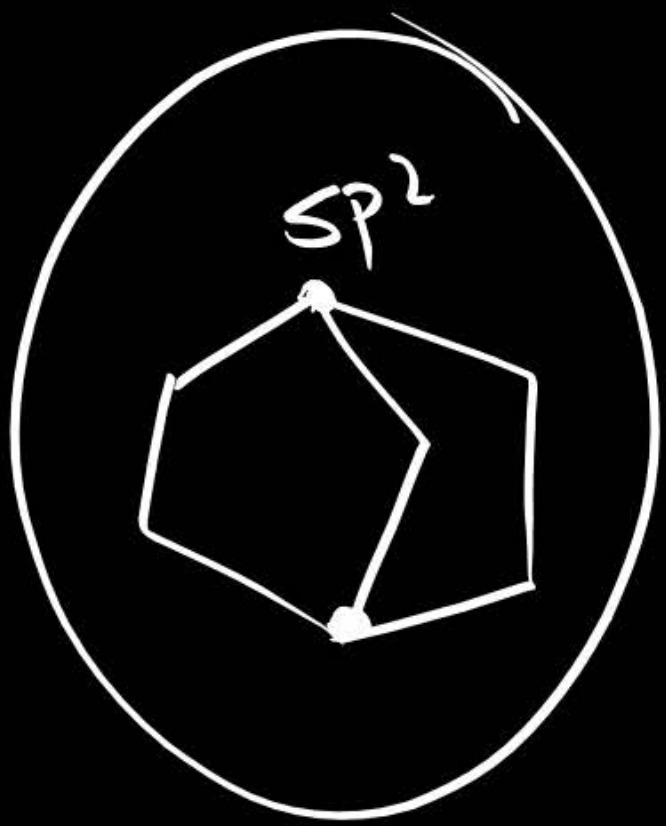
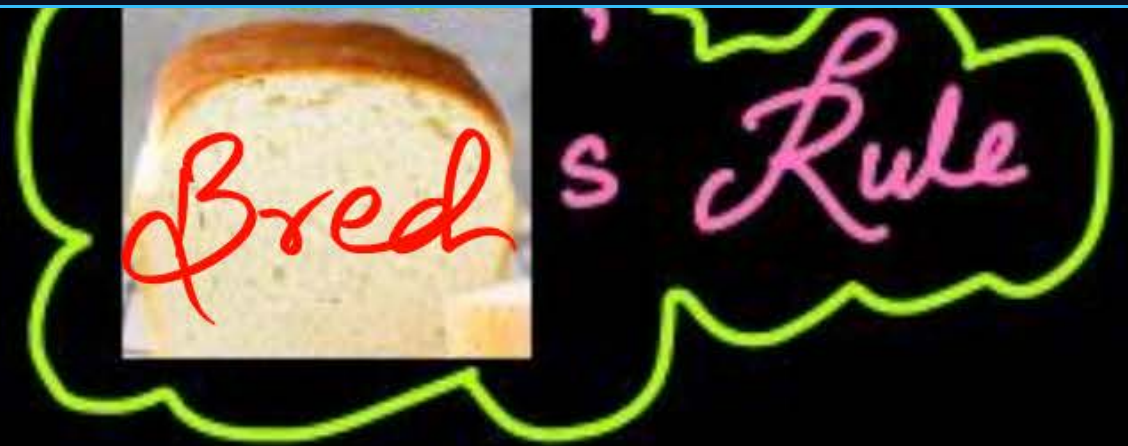
ATDB.uno | 25 Mar 2026, 10:01 IST

ATDB.uno









ring ≤ 8
ATDB.uno
Bridge atom $\Rightarrow$ Planarity (sp^2)

1. Which of the following species or phenomenon are





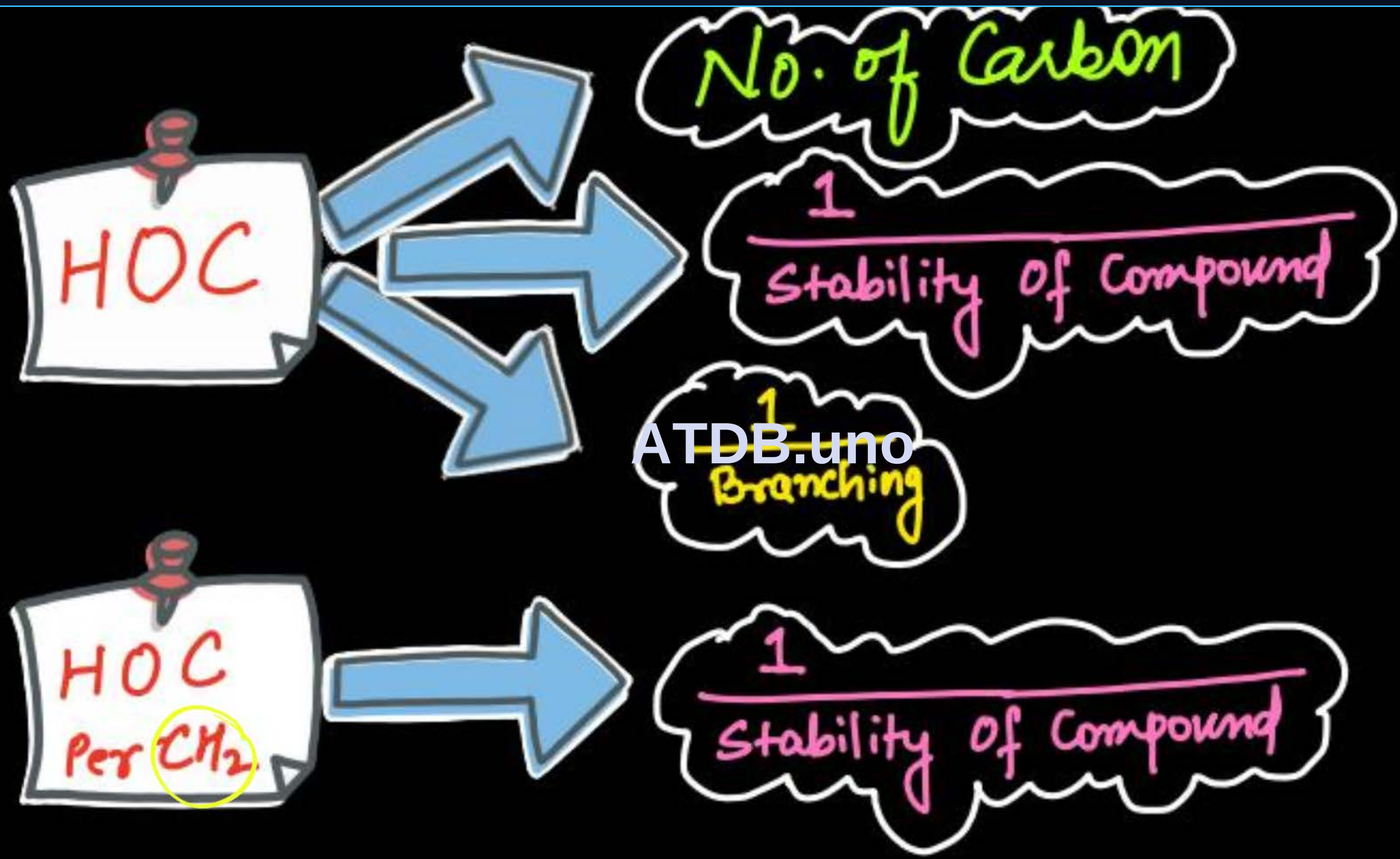
Heat Of Combustion (HOC)

* When one mole of hydrocarbon is fully burned then change in enthalpy is known as 'Heat of Combustion or Enthalpy of Combustion'.

ATDB.uno



$[\Delta H_c]$



Q. Compare HOC (x) and HOC per C (y) for given compounds



①

CCC=C

P

CC=CC

Q

C1=CC=C1

R

(x, y)

Q > R > P

P > R > Q → HOC

②

C12CCC1C2

P

C12CCC1C2

Q

C12CCC1C2

R

Q > R

Q stability → P > Q

③

CC(C)=C

P

CC(C)=C

Q

CC(C)=C

R

CC(C)=C

S

R > P

Q > P → HOC

④

CCC=C

P

CC=CC

Q

6

C1=CC=C1

P

C1=CC=CC1

Q

C1=CC=CC=C1

R

C1=CC=CC=C1

S

HOC per CH₂ → P > Q > R > S

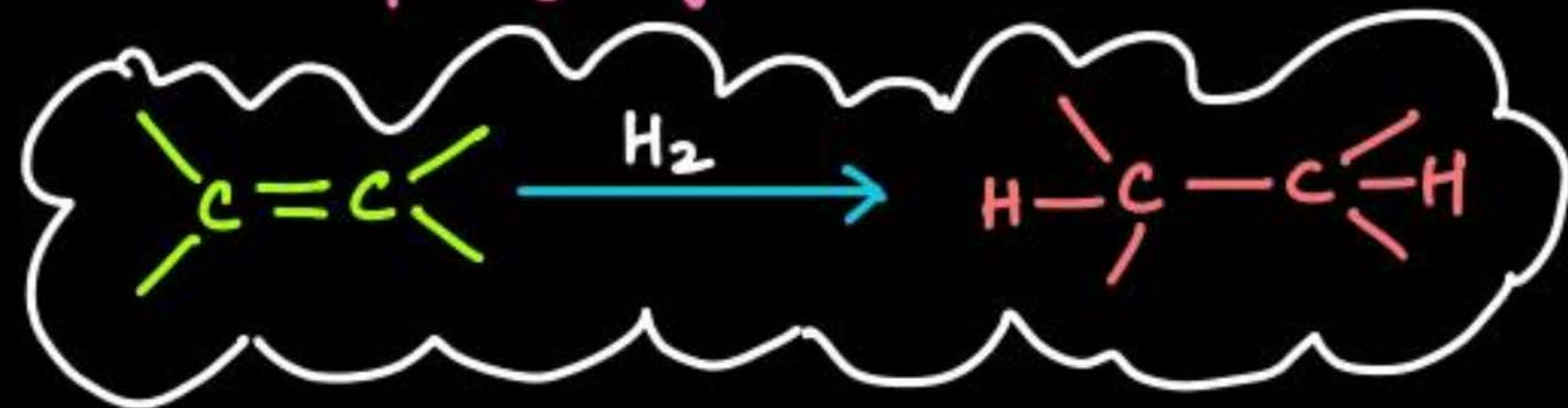
Heat Of Hydrogenation (HOH)



* When 1 mole of a compound completely reacts with 1 mole of Hydrogen(H_2) then change in enthalpy is

Known as 'Heat of Hydrogenation'

Ex.



[ΔH_H]

1.

No. of π -bond

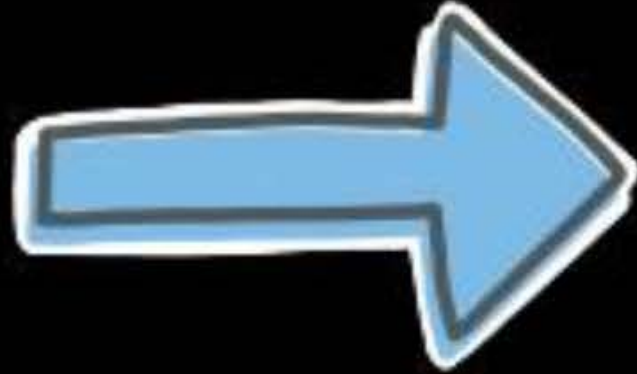


ATDB.uno

1
Stability of Compound *

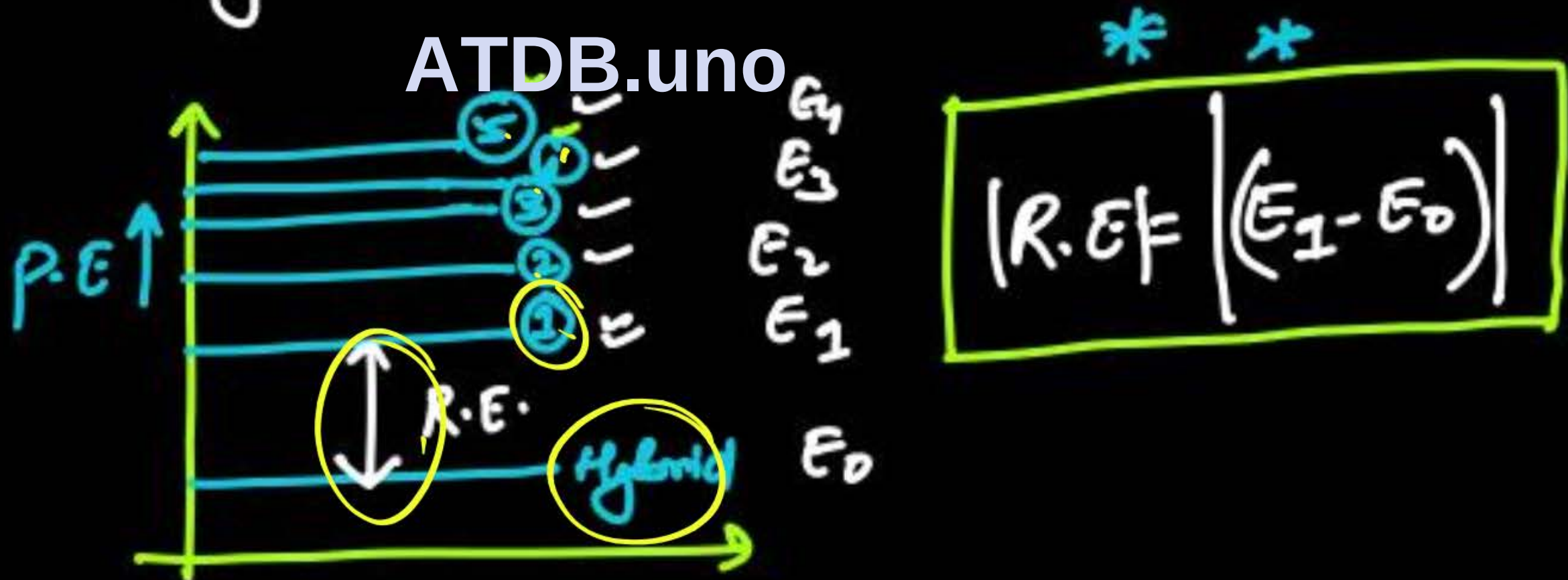
2.

1
Stability of Compound *



Resonance Stabilisation Energy

* Energy difference b/w most stable resonating structure
+ Resonance Hybrid is known as Resonance Energy.





Q. Compare the resonance stabilisation energy for given pair ?





Calculation of Enthalpy



-28.6

$\Delta H_H = -28.6 \text{ Kcal/mol}$



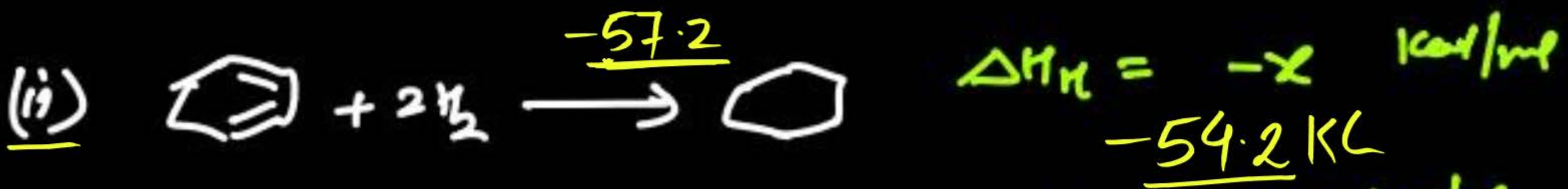
$\Delta H_H = -54.0 \text{ Kcal/mol}$

* find the resonance energy?
 $\Rightarrow -3.2 \text{ Kcal/mol}$

-57.2

-

-3.2

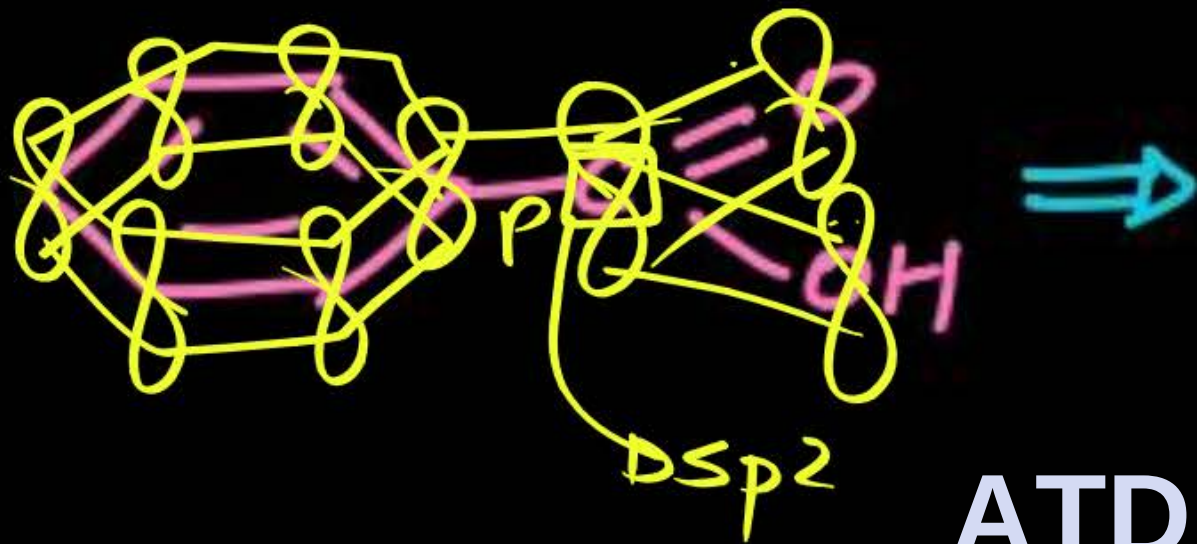


* find the resonance stabilization energy of benzene & also
find ΔH_H for cyclohexadiene if R.E. is -3 kcal/mol ?

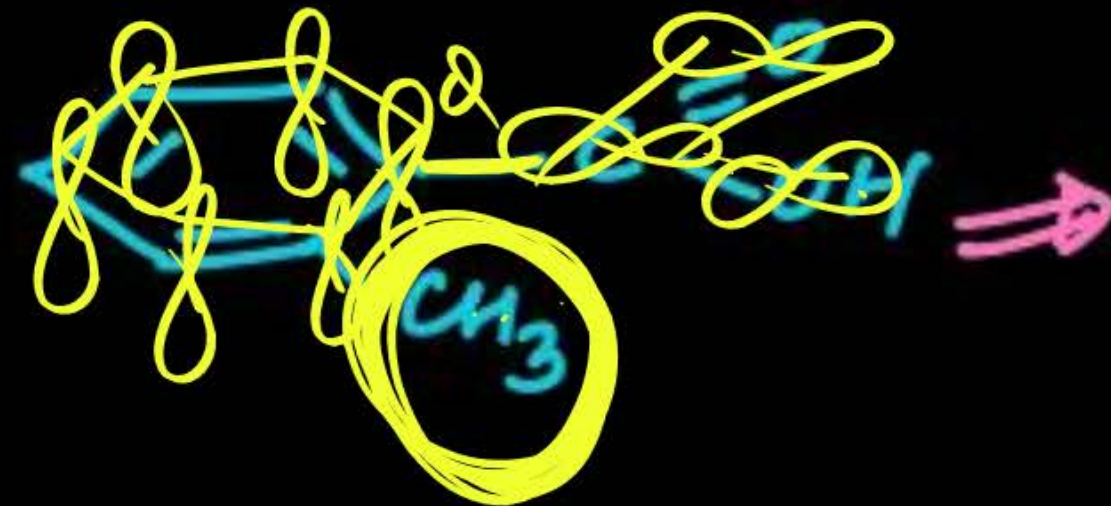


Steric Inhibition Of Resonance (SIR Effect)





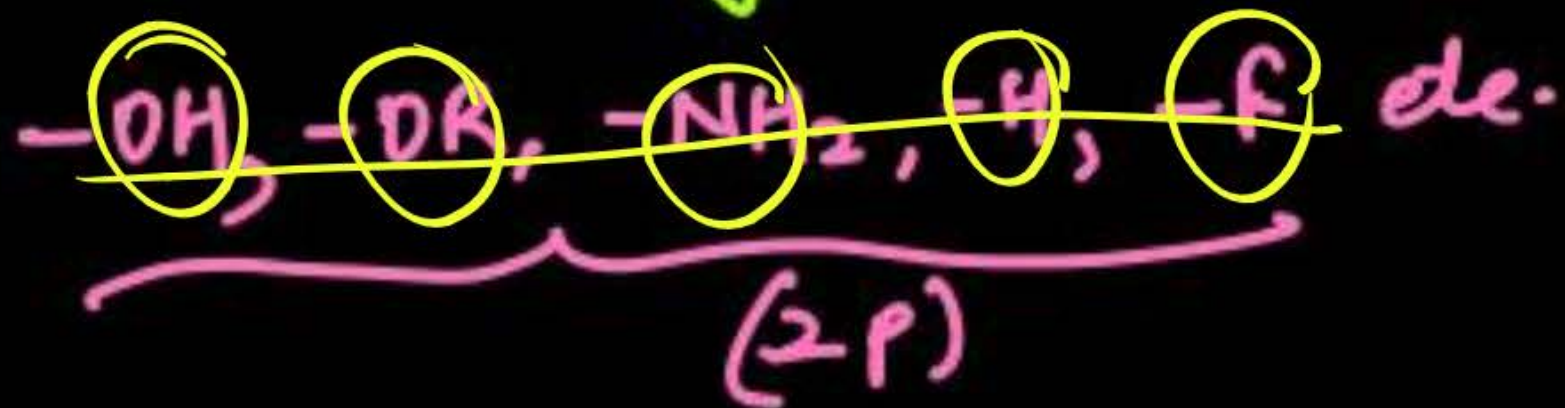
ATDB.uno



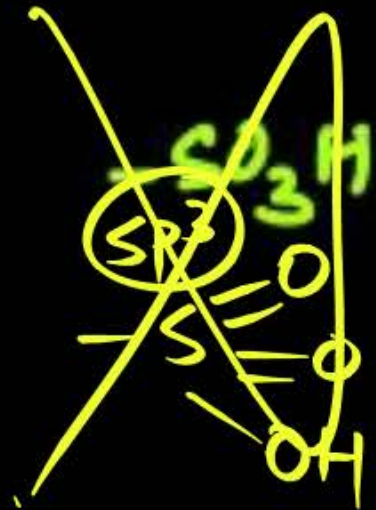
* Due to steric repulsion by group (G) at ortho position, $-sp^2$ group becomes non planar and resonance will inhibit. This inhibition of resonance is known as S.I.R. effect.

ATDB.uno

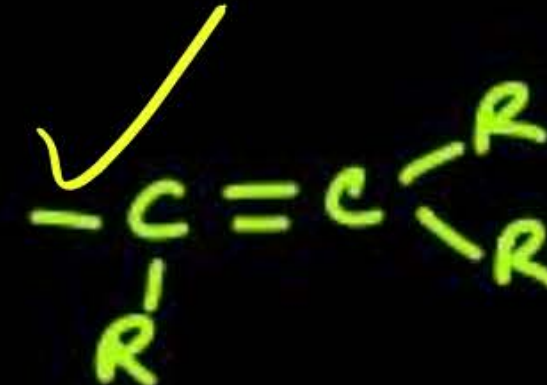
* S-I.R. applied by: All bulky groups except



* SIR applied on: [sp² group]



ATDB.uno



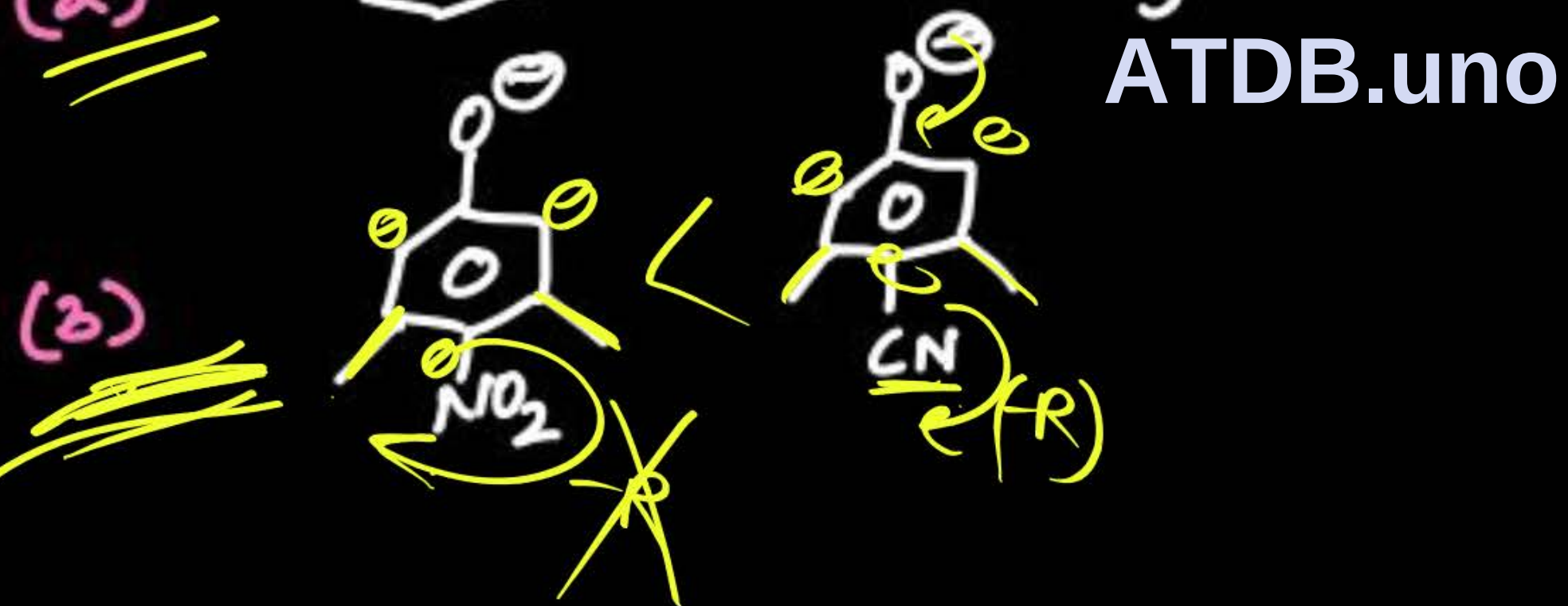
Q. Find the correct order of given properties ?



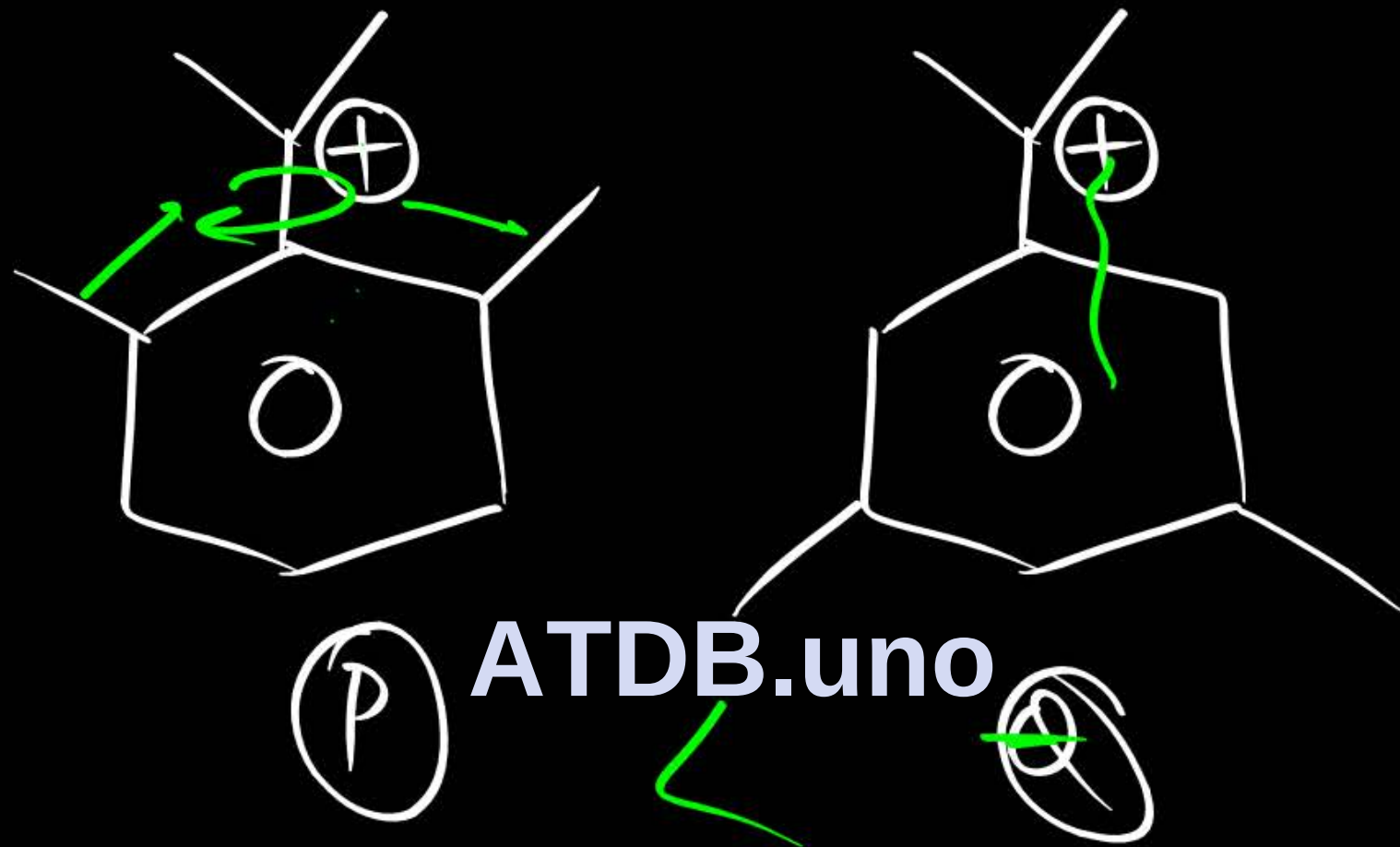
[Bond length]



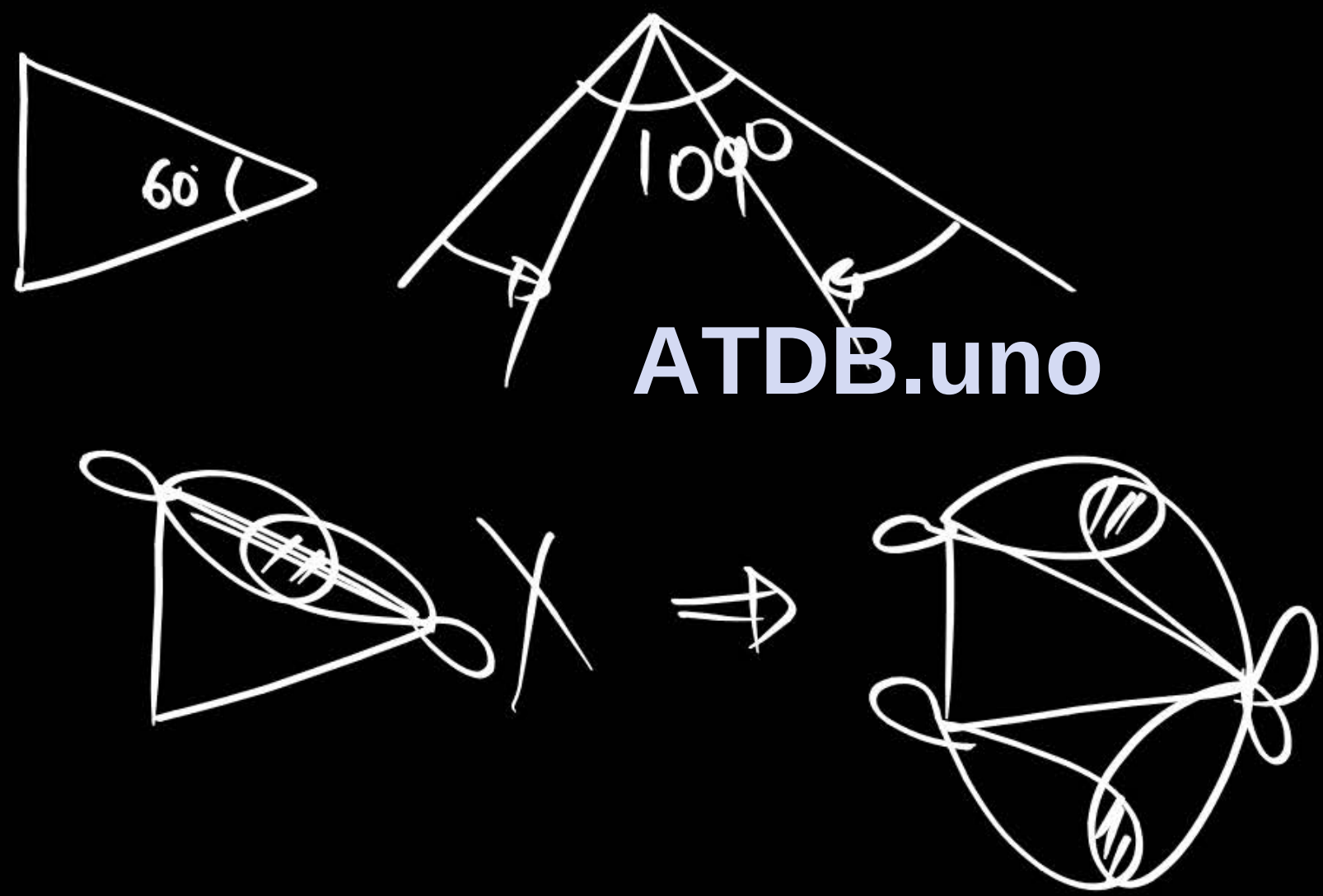
[Resonance Effect]

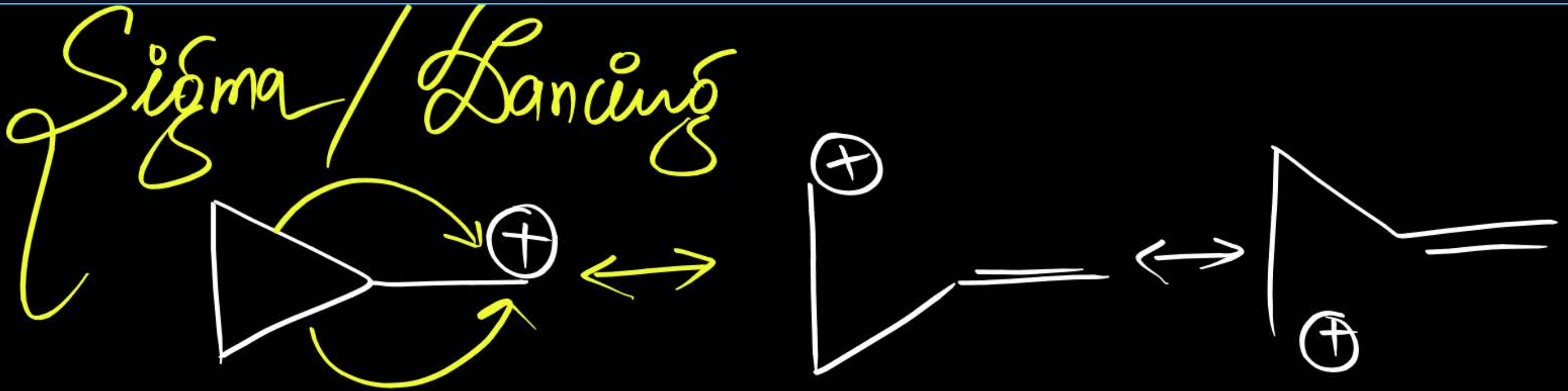


[Stability of (-)ve]



Cyclopropane





ATDB.uno



👉 Bond Dissociation Energy $\Rightarrow$ Homolysis

🎉 The amount of energy required to break a bond is same as the amount of energy released when the same bond is formed.

🎉 In gaseous state, the energy required for homolytic cleavage of a bond is called Bond Dissociation Energy (BDE) or Bond Strength.

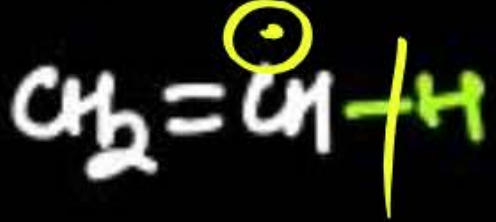
🎉 BDE is affected by s-character of the bond and the stability of the radicals formed. Shorter bonds are typically stronger bonds. $\hookrightarrow$

$$\text{B.D.E.} \propto \frac{1}{\text{Stability of Radical}}$$

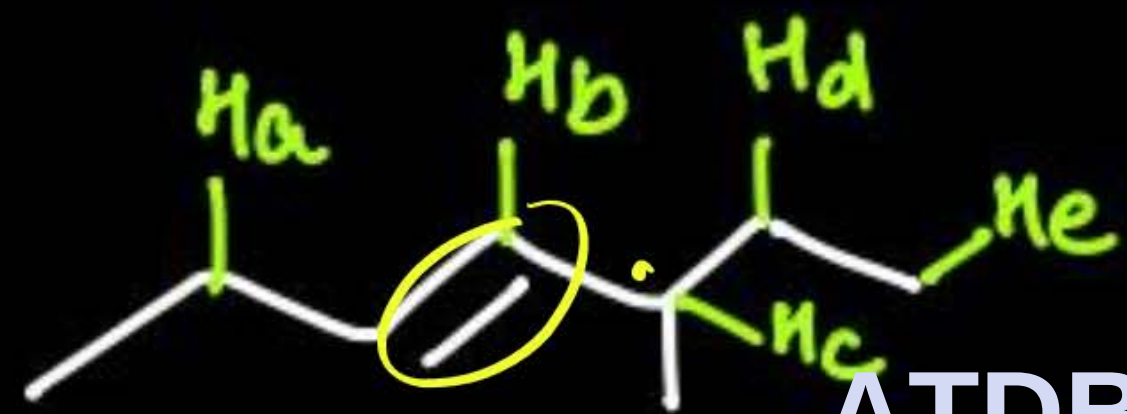
Q. Find the bond energy of given bonds ?



(1)



(2)



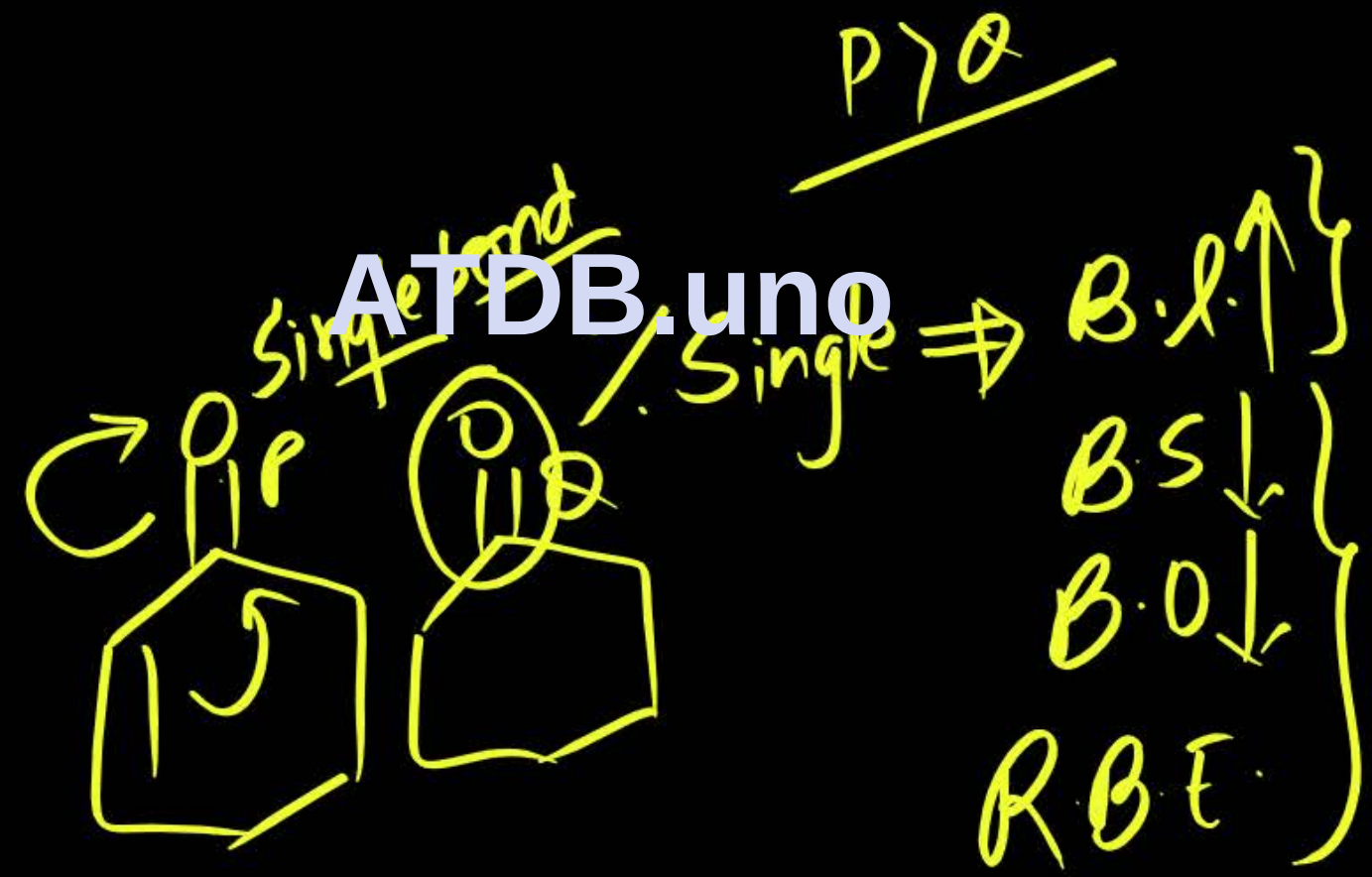
* 1>2>3 => delta + a.
* 3>2>1 => B-D-E.

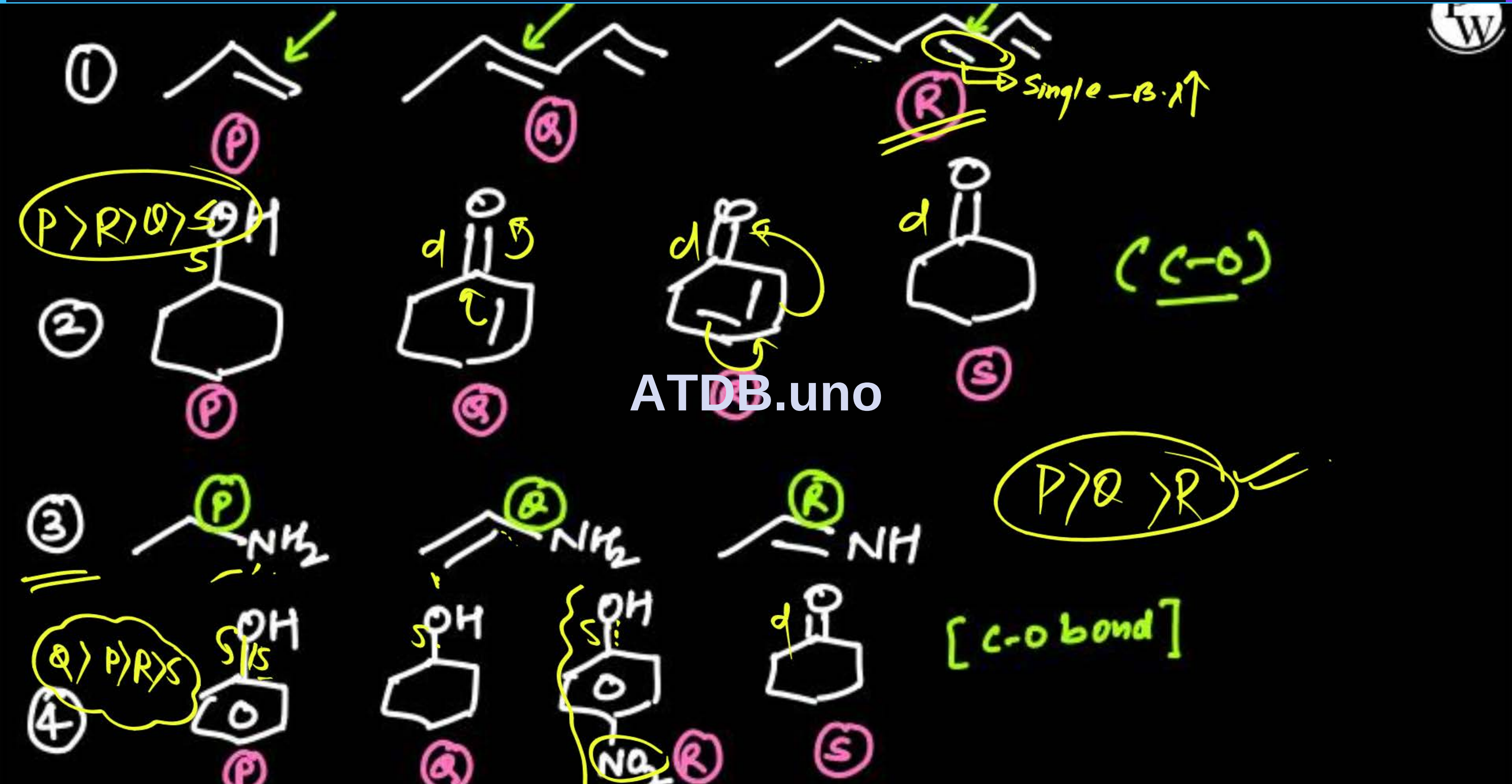
ATDB.uno

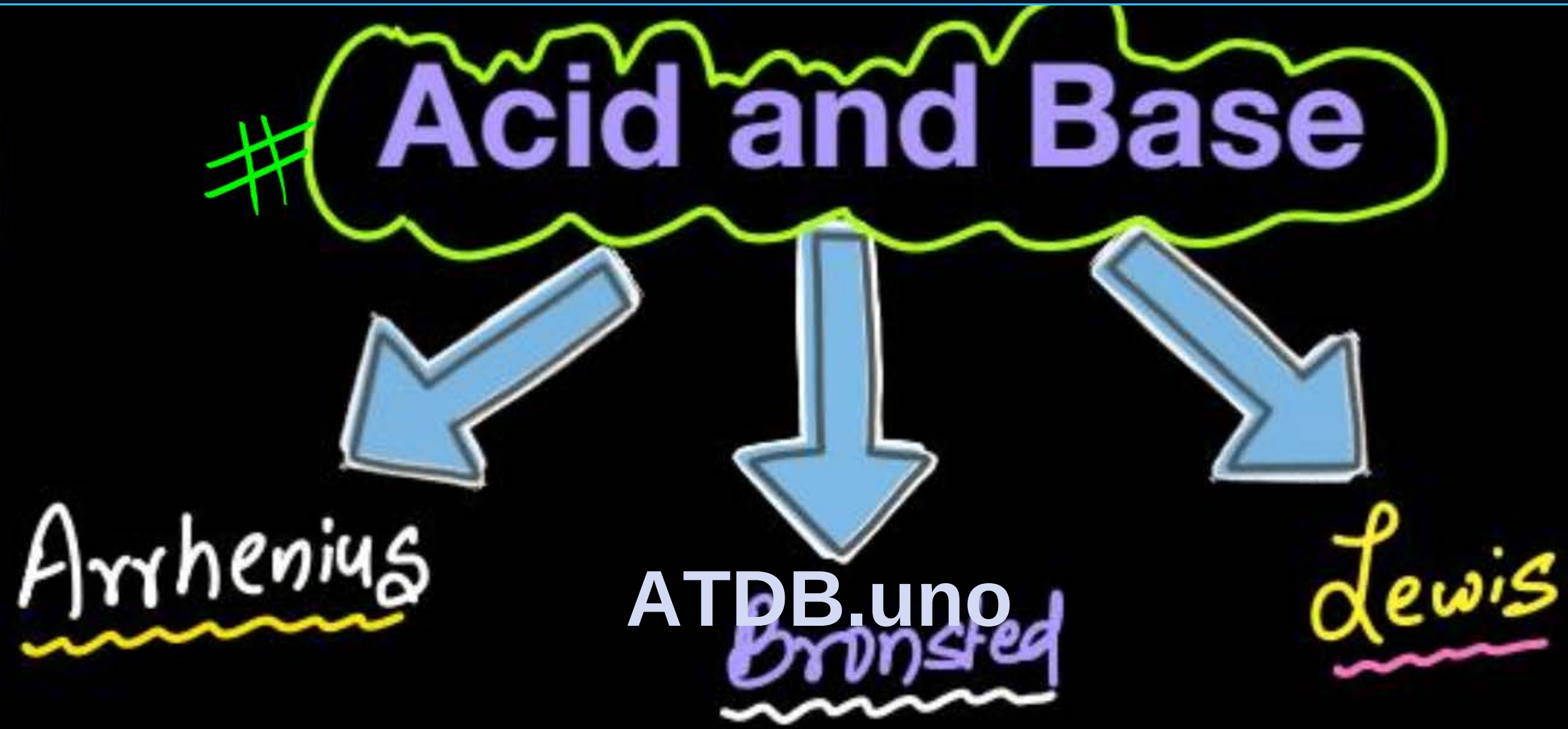
delta + a b i => c > a > d > e > b
||
BDE A = b > e > d > a > c

c a d e b

Bond length

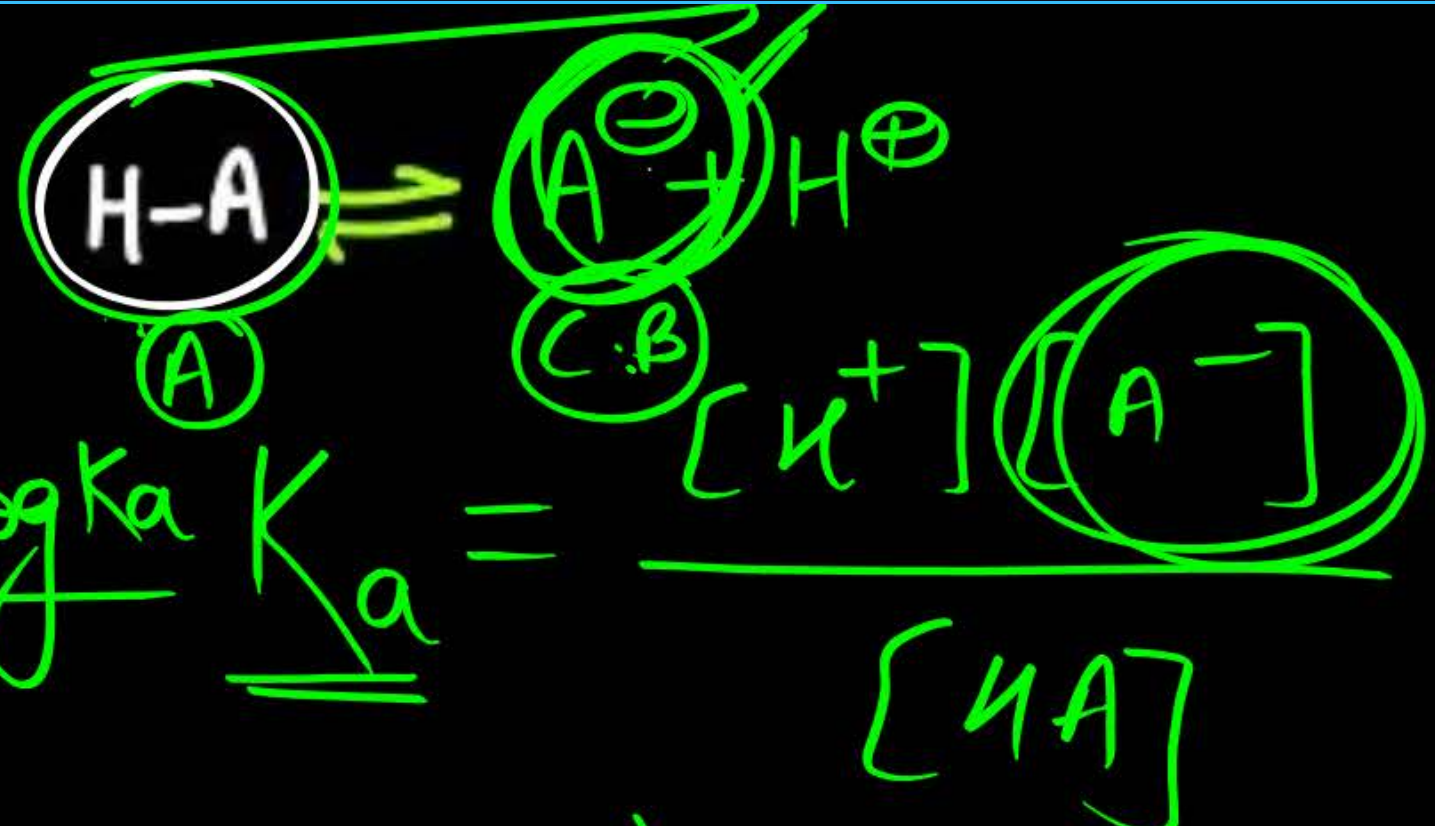






Acidic Strength

EN $\rightarrow$
Size $\downarrow$



- A[⊖] Stability $\Rightarrow$
- ① EWG

② -R, -I (EN)

③ $\frac{1}{+R, +H, +I}$

① H-bonding of C.B. $\uparrow$

② SdR $\uparrow$

③ -R $\uparrow$

④ H-bonding of acid $\downarrow$

⑤ -I



B.T.S.

* Ortho substituted carboxylic acid are most acidic among other isomers (m,p) due to SIR effect.

ATDB.uno

* H-bonding in conjugate base will increase the stability but

if H-bonding only occurs in Acid then acidic strength decreases.



(1) HF (P) HCl (Q) HBr (R) HI (S)

(2) CH4 (P) NH3 (Q) H2O (R) HF (S)

(3)

(3) CH_2OH^- (P), $\text{CH}(\text{OH})^-$ (Q), $\text{C}(\text{OH})(\text{R})^-$ (R)
Below: PhOH^- (P), RCOOH^- (Q), H_2O (R), RSO_3H^- (S)
Central watermark: ATDB.uno

(4)

(4) CH_2OH^- (P), $\text{CH}(\text{OH})^-$ (Q)
Below: cyclopropyl-OH (Q), cyclobutyl-OH (Q), cyclopentyl-COOH (R), cyclohexyl-COOH (S)
Central watermark: ATDB.uno

(5)

(5) PhOH^- (P), RCOOH^- (Q)
Below: H_2O (R), RSO_3H^- (S)
Central watermark: ATDB.uno

(6)

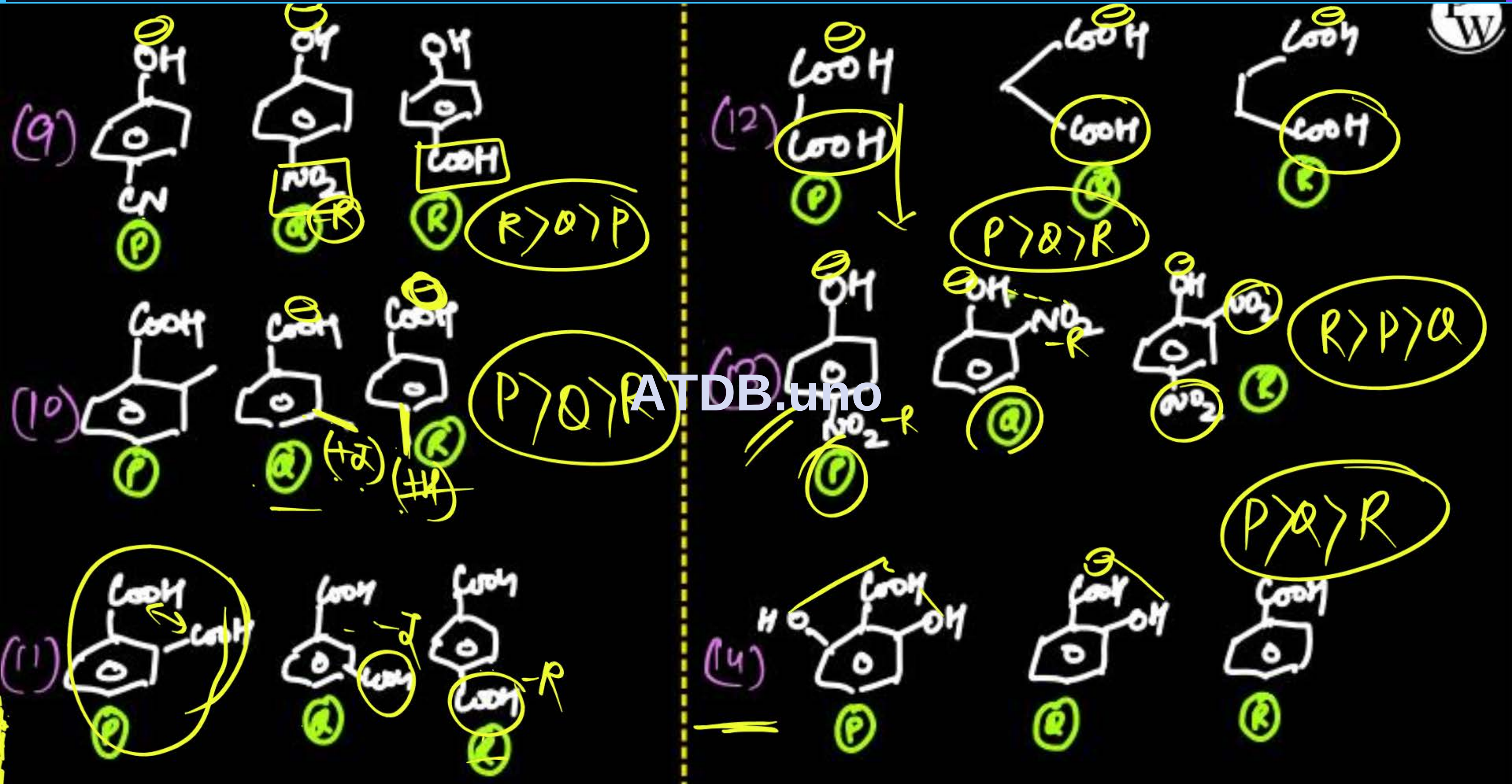
(6) CH_2OH^- (P), $\text{CH}(\text{OH})^-$ (Q)
Below: cyclopropyl-OH (Q), cyclobutyl-OH (Q), cyclopentyl-COOH (R), cyclohexyl-COOH (S)
Central watermark: ATDB.uno

(7)

(7) $\text{HC}\equiv\text{CH}$ (P), $\text{CH}_2=\text{CH}_2$ (Q), $\text{CH}_2-\text{C}\equiv\text{CH}$ (R)
Below: HCl (S), H_2O (R), HCl (S)
Central watermark: ATDB.uno

(8)

(8) $\text{HC}\equiv\text{CH}$ (P), NH_3 (Q)
Below: H_2O (R), HCl (S)
Central watermark: ATDB.uno



Basic Strength



$B^{\ominus} \Rightarrow$ ① $e^{\ominus}$ donating tendency

ATDB.uno

Localised

✓ (b) + R

(c) + H

(d) + I

use



B.T.S.

* Ortho substituted aniline are least basic among other isomer. [Solvation Effect]

ATDB.uno

* As the electron density at donor site increases by +R, +M, +I, Basic strength increases.



Solvation Effect (SIP)



#

$\text{-COOH} \Rightarrow \text{AS} \uparrow$

$\text{-NH}_2 \Rightarrow \text{BS} \downarrow$

ATDB.uno

①

$F^- > Cl^- > Br^- > I^-$

③

$H_2O < NH_3$

⑤

$R-OH < R-NH_2$

⑦

$CH_3NH_2 < CH_2=NH < HC \equiv N$

⑨

$Ph-NH_2 < Ph-CH_2NH_2 < Ph-NH-CH_3 < Ph-N(CH_3)_2$

②

$CH_3^- < NH_2^- < OH^- < F^-$

④

$NaOH < Na < NaNH_2$

⑥

$R-COO^- < R-O^- < Ph-O^-$

⑧

$Ph-NH_2 < Ph-NHMe < Ph-NMe_2$

⑩

$Ph-NH_2 < Ph-CO-NH_2 < Ph-NH-CO-CH_3$

②

I^-

③

NH_2^-

⑦

$HC \equiv N$

⑨

$Ph-NH-CH_3$

⑩

$Ph-N(CH_3)_2$

Size

EN ↑

$P > O > R$

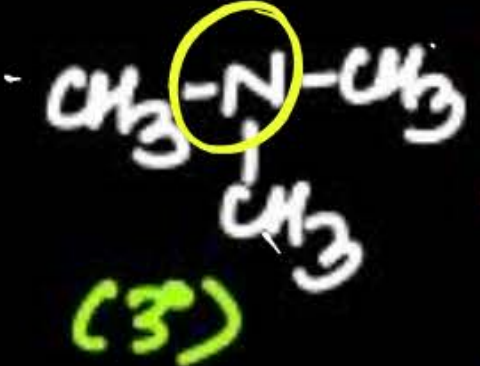
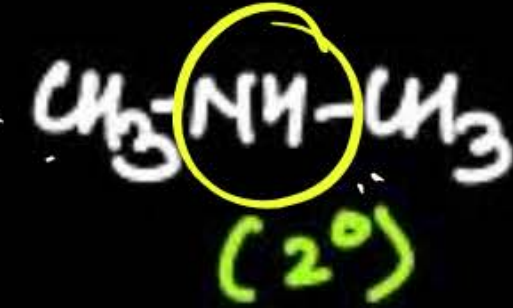
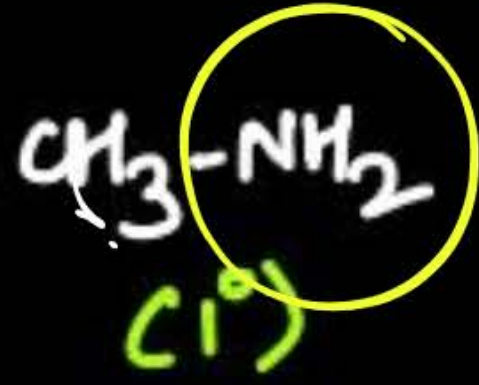
$N > R > P > S$

ATDB.uno





Basic Strength In Water



Gas phase
3 > 2 > 1

Aqueous phase

1° 3°

213
231

2 ⇒ ATDB.uno

H-bond ⇒ 1 > 2 > 3

* 2 > 1 > 3 ⇒ Me
* 2 > 3 > 1 ⇒ Et

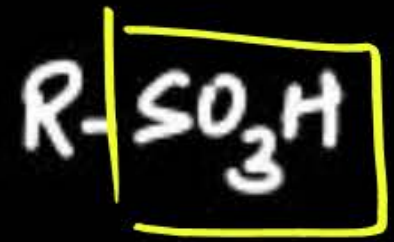
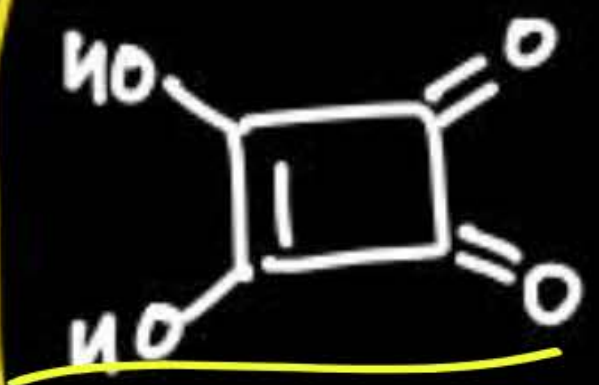
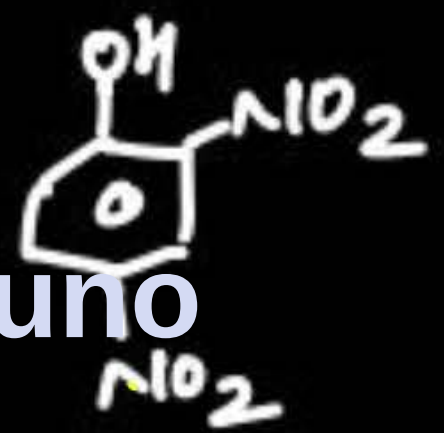
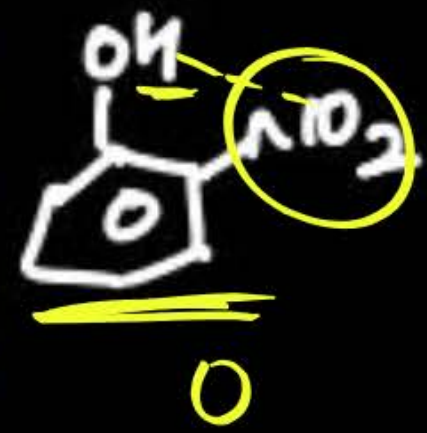
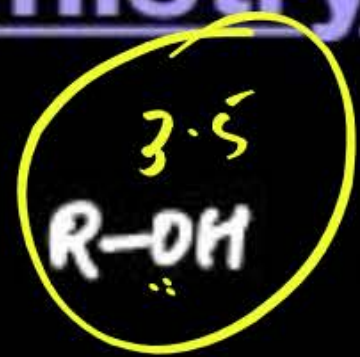


Solubility Table

Practical Organic Chemistry



Alkane Alkene



Inorganic acids

($HNO_3, H_2SO_4, HCl, HBr, \dots$)



Solubility in Aq. NaOH



* If HA is more acidic than H_2O then reaction will move in forward direction.

* If reaction moves in forward direction then HA will be Soluble in aq. NaOH.



Solubility in Aq. NaHCO_3



* If HA is more acidic than H_2CO_3 then reaction will move in forward direction.

* If reaction moves in forward direction then HA will be Soluble in aq. NaHCO_3 .



example

Differential
Extraction



ether
aq. NaOH

PhO ⁻	H ₂ O
<chem>c1ccccc1</chem>	ether

3



Organic Solvent [Ether]



Aq. HCl
acid
PhNH₃⁺

(Then aq. NaOH)



Ph-NH₂ Q

Aq. NaHCO₃

(Then aq. HCl)



PhCOOH

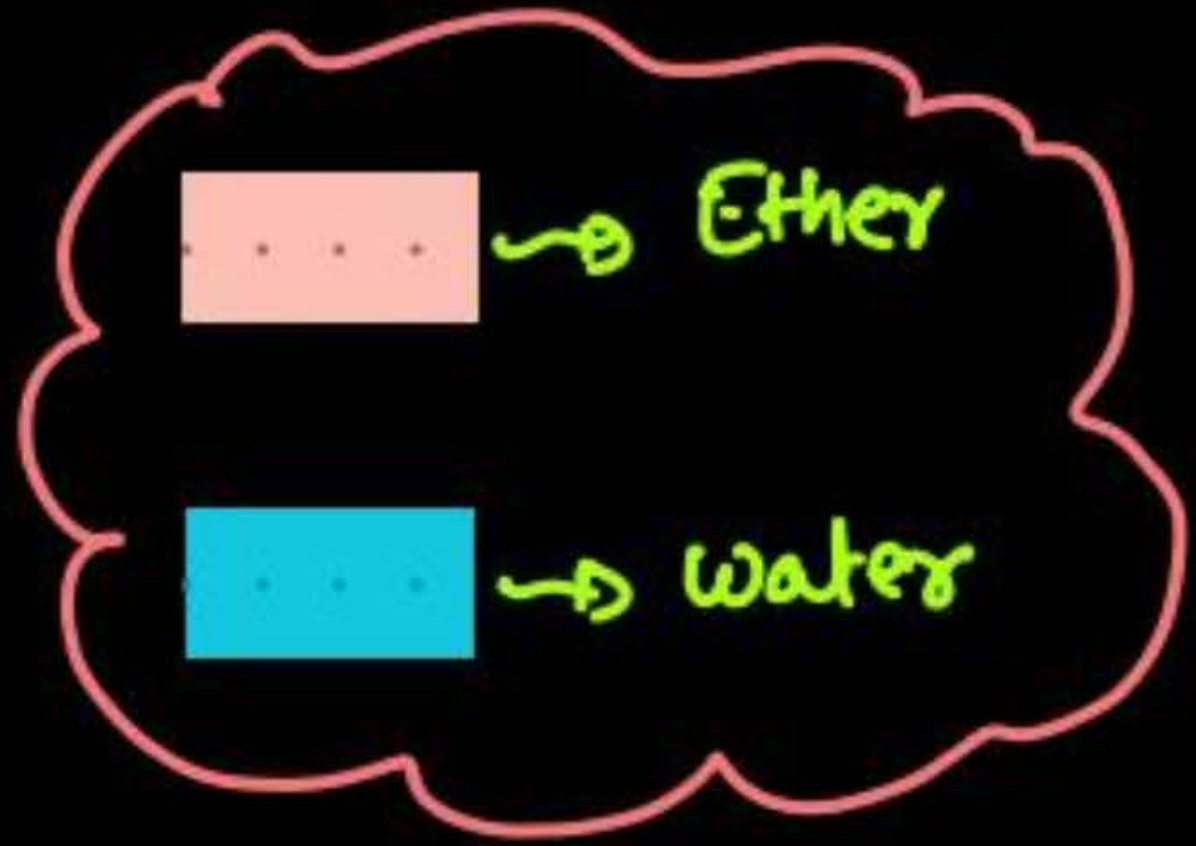


Aq. NaOH

(Then aq. HCl)



PhOH



ATDB.uno

