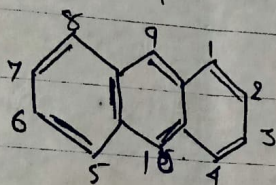


Anthracene -

Extraction from Coal tar $\rightarrow$ Anthracene occurs in Coal tar (less than 1%) and is obtained from green oil fraction (b.p. $270^\circ\text{C} - 366^\circ\text{C}$). On cooling this fraction crude Anthracene crystallised out. Crude Anthracene contains Phenanthrene and Carbazole as impurities.

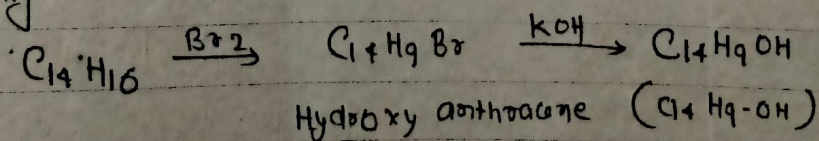
It is purified by washing it with solvent naphthalene (to remove Phenanthrene) and Pyridine (to remove Carbazole). Finally the solid is sublimation to give pure Anthracene.

In naming derivatives of Anthracene, the numbering system shown below is used.

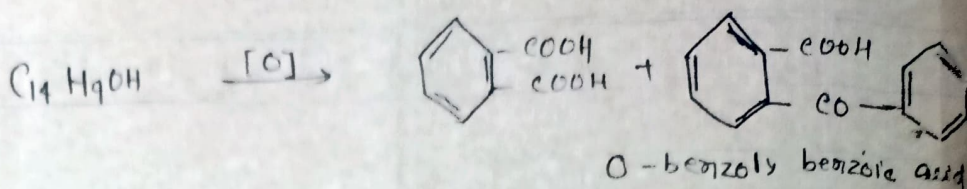


Classical Structure of Anthracene $\rightarrow$

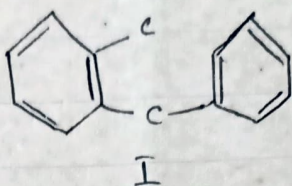
- (1) It has a molecular formula $\text{C}_{14}\text{H}_{10}$
- (2) It undergoes the typical aromatic substitution reaction like Sulphonation, Nitration, halogenation etc. It shows the aromatic character of Anthracene.
- (3) Bromination of Anthracene gives 9-bromo Anthracene, which on fusion with KOH gives hydroxy Anthracene.



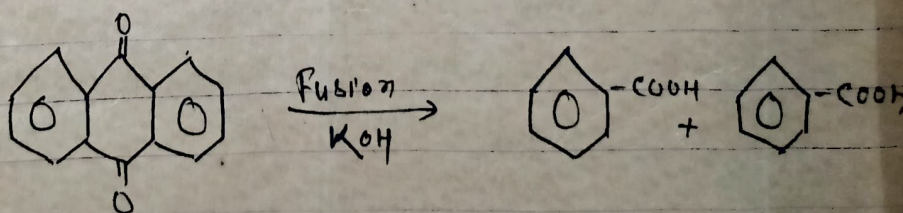
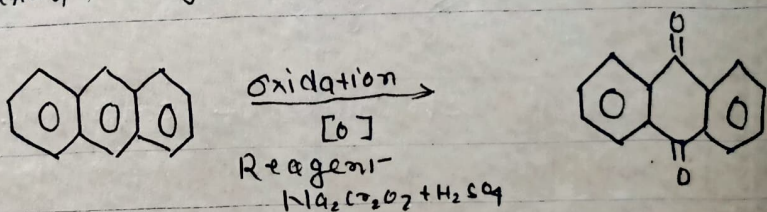
On vigorous oxidation gives Phthalic acid and a small amount of o-benzoyl benzoic acid.



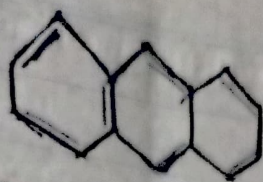
This suggests that anthracene contains at least two benzene rings and that its skeleton is as shown.



The presence of two benzene ring is confirmed by the fact that on fusion with KOH at $250^{\circ}C$, anthraquinone (which may be obtained from anthracene by direct oxidation) gives two molecules of benzoic acid.

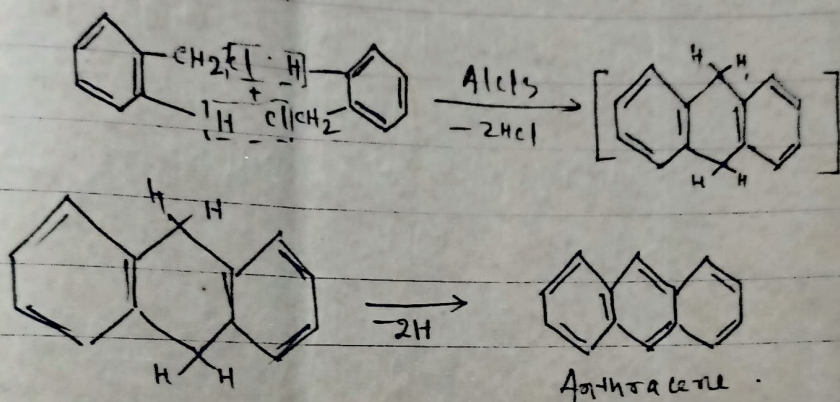


The above reaction confirmed the presence of two benzene rings in anthracene. In skeleton I, there are 14 C-atoms. Now, to fit 10 H atoms and to retain the quadrivalency of C-atoms, the middle ring should be closed. On taking these facts into consideration following structure of anthracene have been suggested.

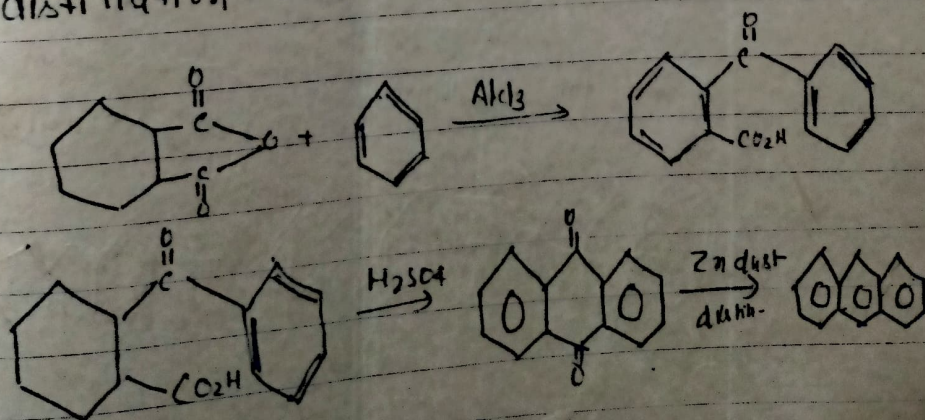


Above Structure has been confirmed by following Synthesis $\rightarrow$

(1) By Fiedel-Craft - Reaction $\rightarrow$ Benzyl chloride reacts with it self to form 9,10-dihydroanthracene which readily loses two H-atoms to yield anthracene.

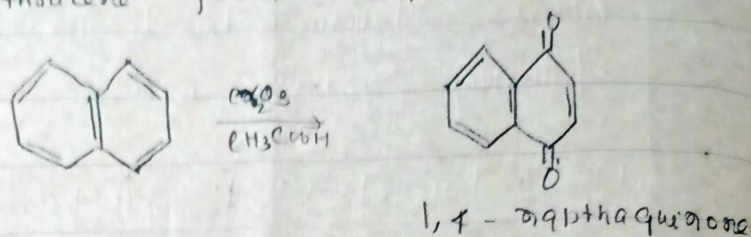


(2) Haworth Synthesis $\rightarrow$ When Phthalic anhydride in benzene solution is treated with AlCl_3 , o-benzoyl Benzoic acid is formed. This on heating with Conc. H_2SO_4 at 100°C forms 9,10-anthraquinone, which on distillation with Zn dust gives anthracene.

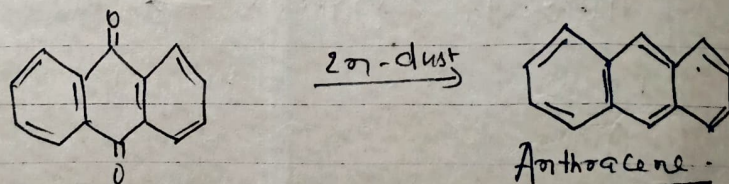
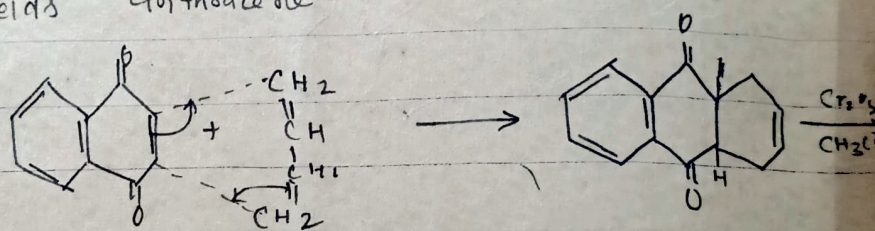


(B) By Diels Alder reaction $\rightarrow$

This reaction is used to synthesizing anthracene from naphthalene.



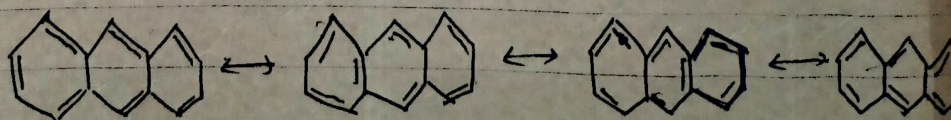
1,4-naphthoquinone undergoes Diel-Alder-reaction with 1,3-butadiene. The product of this reaction is oxidised with Cr_2O_3 in glacial acetic acid to form, 9,10-anthraquinone, Distillation of anthraquinone with Zn-dust yields Anthracene.



Anthraquinone.

Resonance hybrid Structure $\rightarrow$

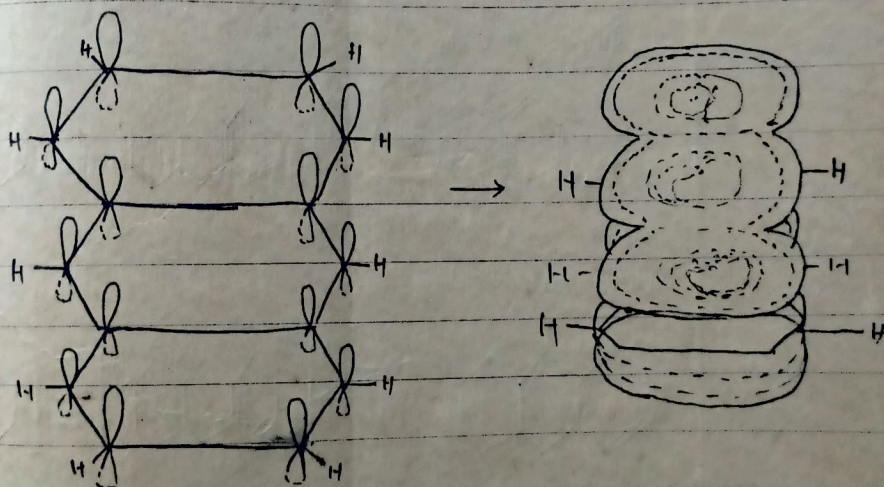
According to resonance theory anthracene is considered to be the resonance hybrid of the following four Canonical forms.



Orbital Picture of Anthracene $\rightarrow$

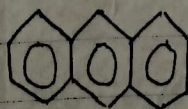
All 14 C-atoms in anthracene are sp^2 hybridized. The sp^2 hybrid orbitals overlap with each other and with s-orbitals of the ten H-atoms forming C-C and C-H bonds. Since the bonds result from the overlap of tetragonal sp^2 orbitals, so all the C-bonds are in one plane.

Each C-atom in anthracene possesses an unhybridised p-orbital perpendicular to the plane of C-bonds. The lateral overlap of these p-orbitals produces a π M.O. containing 14 electrons one half ~~for~~ half of this π M.O. lies above the plane of bonds and other half lies below the plane of C bonds.



Nowadays Anthracene is represented.

as.



Properties of anthracene $\rightarrow$ Anthracene is
 Colourless solid. It melts at 218°C and boils
 at 340°C . It is insoluble in water but
 dissolves in benzene. It shows a strong
 fluorescence when exposed to u.v. light.

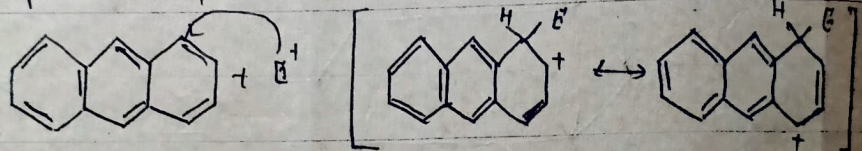
Chemical Properties $\rightarrow$

Anthracene undergoes addition and
 electrophilic substitution reactions. These
 reactions preferentially occur at C-9

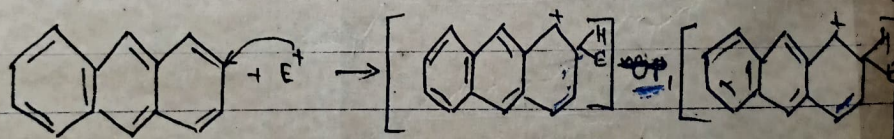
and C-10 positions. This can be understood
 if we examine the intermediate carbocation
 ion obtained from attack at C-1, C-2 and C-9

E^+ in the following equations
 represent an electrophile.

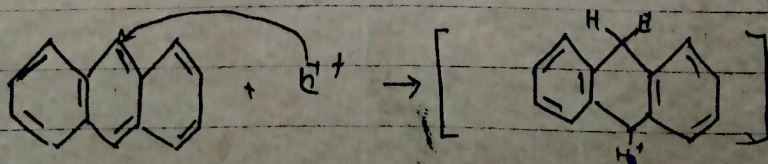
Attack at C-1



Attack at C-2



Attack at C-9



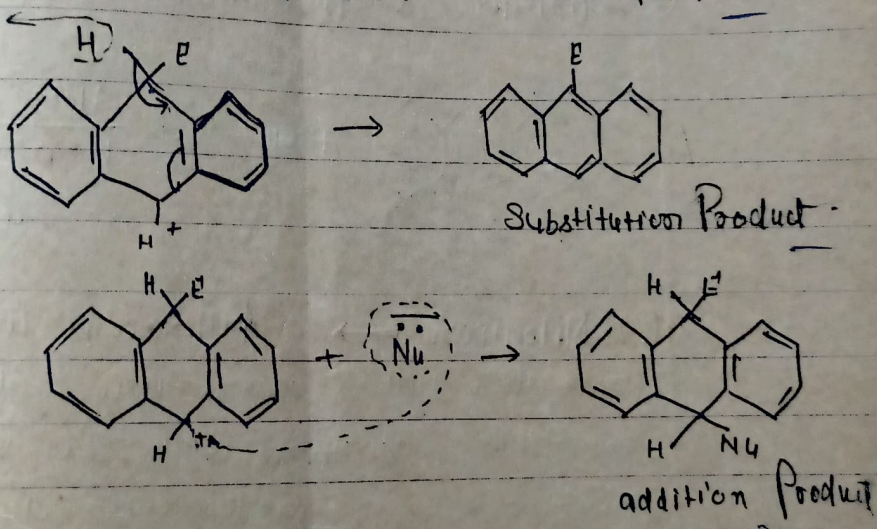
Attack at C-9 $\rightarrow$

Carbocation ion in which two benzenoid rings are retained.

C-1 or C-2 yields an intermediate in which a naphthalene system is retained.

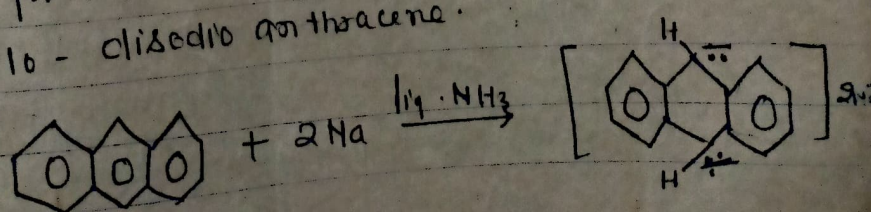
The former intermediate is more stable and its formation favoured because resonance energy ($2 \times 36 = 72 \text{ Kcal}$) of two Benzene rings exceeds that of naphthalene (61 Kcal).

This Carbocation ion can lose a proton to give the corresponding substitution product, or it can react with a nucleophile to form a 9-10 addition product.

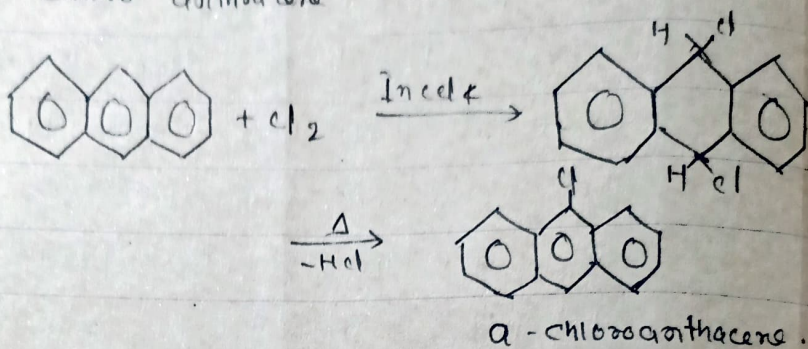


The main chemical Properties of anthracene are described below $\rightarrow$

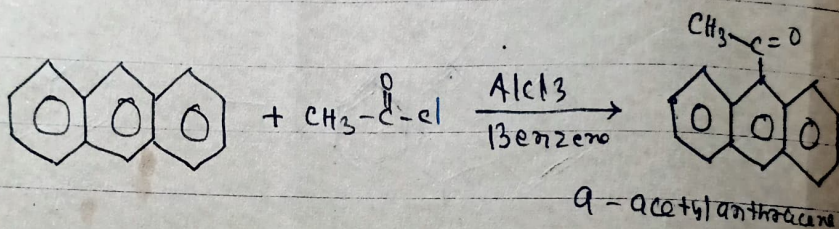
(i) Reaction with Sodium $\rightarrow$ Anthracene reacts with metallic sodium in liquid ammonia to form a deep blue solution of 9,10-disedide anthracene.



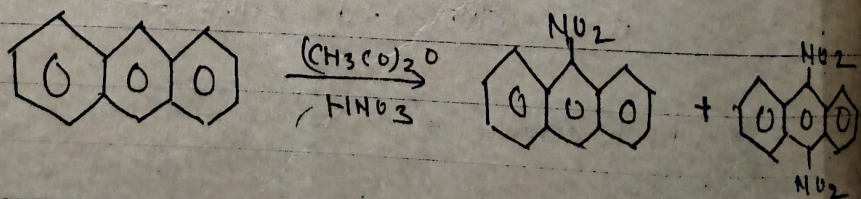
(2) Reaction with halogens $\rightarrow$ Anthracene reacts with Cl_2 in CCl_4 at room temp. to give 9,10-dichloro-9,10-dihydroxyanthracene. On heating this addition product, it loses a molecule of HCl forming 9-chloroanthracene.



(3) Friedel-Crafts acylation $\rightarrow$



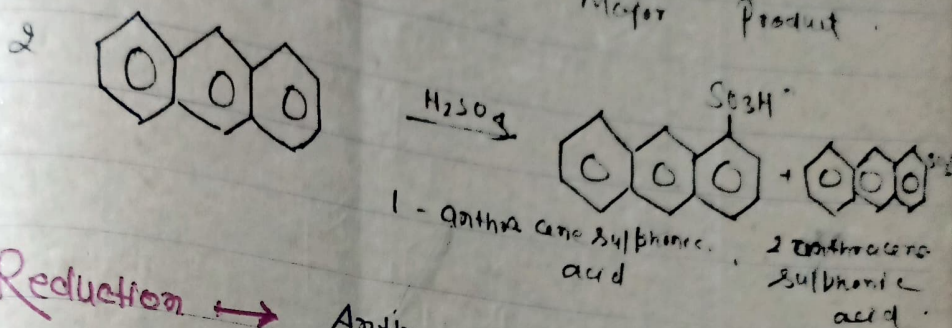
(4) Nitration $\rightarrow$ Anthracene undergoes nitration with conc. HNO_3 in acetic anhydride at room temp. to yield a mixture of 9-nitroanthracene and 9,10-dinitroanthracene.



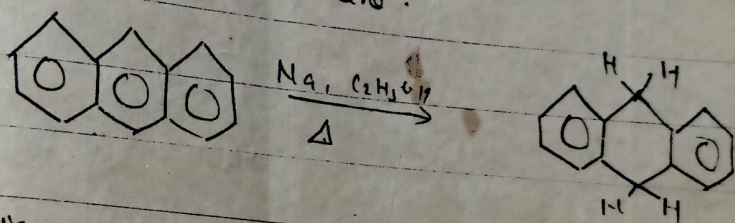
The usual nitrating mixture ($\text{HNO}_3 + \text{H}_2\text{SO}_4$) is not used because it leads to the formation of 9,10-anthraquinone by oxidation.

(5) **Sulphonation** $\rightarrow$ Anthracene undergoes Sulphonation with conc. H_2SO_4 to yield a mixture of 1-anthracene sulphonic acid and 2-anthracene sulphonic acid.

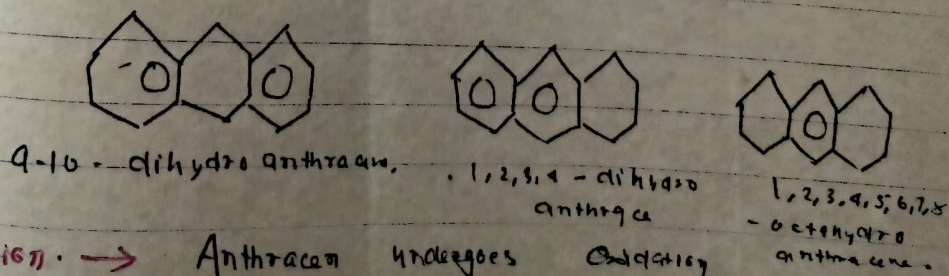
At a lower temperature former is the major product and at a higher temperature latter is the major product.



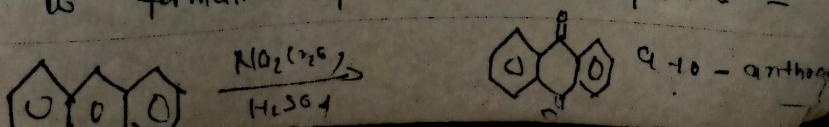
Reduction $\rightarrow$ Anthracene undergoes reduction with sodium and ethyl alcohol to form 9,10-dihydro anthracene.



Catalytic reduction using nickel at $225^\circ C$ first gives 9,10-dihydroanthracene, and by ~~continued~~ continued hydrogenation this is converted into 1,2,3,4-tetrahydroanthracene and 1,2,3,4,5,6,7,8-octahydroanthracene.

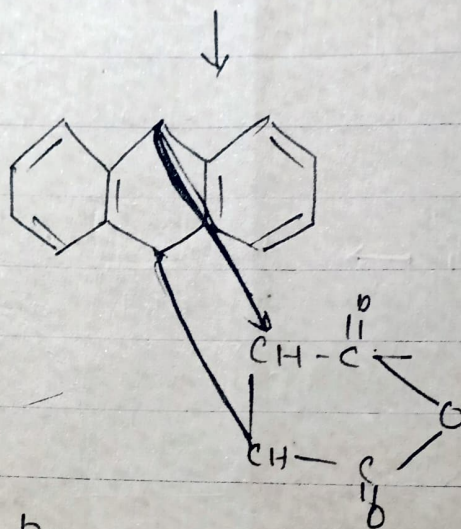
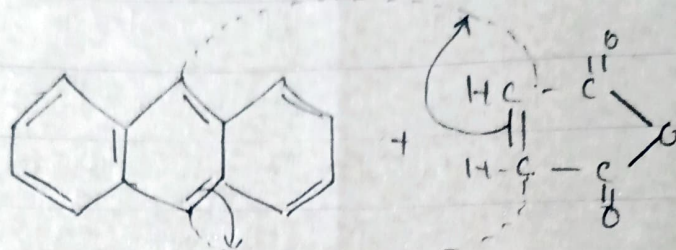


Oxidation $\rightarrow$ Anthracene undergoes oxidation with $(Na_2Cr_2O_7 + H_2SO_4)$ to form 9,10-anthraquinone. Other oxidizing agent like HNO_3 also V_2O_5 also led to formation of 9,10-anthraquinone.



Diels - Alder - Reaction $\rightarrow$ Anthracene.

Undergoes a Diels Alder reaction with maleic anhydride to yield the corresponding adduct.



Diels - Alder adduct -

Anthracene is used in the manufacture of anthraquinone.

Ans -