

A-Level Chemistry

Name: _____ Score: _____

1. Benzene reacts mostly by electrophilic substitution because:
 - ☐ this keeps the stable delocalised ring, while addition would destroy it
 - ☐ the ring is unstable
 - ☐ it has no electrons
 - ☐ addition is impossible
2. A halogen reacts in the ring (not the side-chain) when there is:
 - ☐ a halogen-carrier catalyst (AlCl_3) and no UV light
 - ☐ UV light and no catalyst
 - ☐ only heat
 - ☐ water present
3. Nitration of benzene uses:
 - ☐ concentrated HNO_3 and concentrated H_2SO_4
 - ☐ bromine water
 - ☐ dilute NaOH
 - ☐ hydrogen and a catalyst
4. Under UV light with no catalyst, the halogen substitutes:
 - ☐ in the side-chain (free-radical substitution)
 - ☐ in the ring
 - ☐ nowhere
 - ☐ at the catalyst
5. A Friedel–Crafts reaction (with AlCl_3) is used to:
 - ☐ add an alkyl or acyl group to the ring
 - ☐ remove a hydrogen as a radical
 - ☐ oxidise the ring
 - ☐ hydrogenate the ring
6. Groups already on a benzene ring direct where the next group attaches.
 - ☐ True ☐ False

7. Benzene reacts mainly by electrophilic substitution, not addition, because that keeps its stable delocalised ring intact.
- ☐ True ☐ False
8. An -OH group already on the ring directs the next substituent to:
- ☐ positions 2 and 4
- ☐ position 3 only
- ☐ no particular position
- ☐ the side-chain
9. In electrophilic substitution, after the ring bonds to the electrophile:
- ☐ an H^+ is lost, restoring the stable ring
- ☐ the ring opens up
- ☐ another electrophile adds
- ☐ a radical forms
10. Arenes such as benzene typically react by:
- ☐ electrophilic substitution
- ☐ electrophilic addition
- ☐ nucleophilic substitution
- ☐ free-radical addition

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1. **this keeps the stable delocalised ring, while addition would destroy it**

Substitution preserves the aromatic stabilisation; addition would break up the delocalised system.

2. **a halogen-carrier catalyst (AlCl_3) and no UV light**

The catalyst makes the electrophile for ring substitution; UV with no catalyst gives side-chain free-radical substitution.

3. **concentrated HNO_3 and concentrated H_2SO_4**

The conc. acid mixture generates the NO_2^+ electrophile, giving nitrobenzene.

4. **in the side-chain (free-radical substitution)**

UV light starts free-radical substitution in the alkyl side-chain.

5. **add an alkyl or acyl group to the ring**

Friedel-Crafts alkylation/acylation adds a carbon-based group to benzene.

6. **True**

This is the directing effect.

7. **True**

Addition would break the delocalised ring and lose its stability, so substitution (swapping an H) is favoured.

8. **positions 2 and 4**

$-\text{NH}_2$, $-\text{OH}$ and $-\text{R}$ are 2,4-directing; $-\text{NO}_2$, $-\text{COOH}$ and $-\text{COR}$ are 3-directing.

9. **an H^+ is lost, restoring the stable ring**

Losing H^+ from that carbon restores the delocalised ring, completing the substitution.

10. **electrophilic substitution**

Substitution keeps the stable ring intact.