F324: Rings, Polymers and Analysis 4.1.4 Amines / 74

1.

$$H_3C$$
 H_3C
 $\uparrow N \equiv N$

ALLOW ECF ✓✓ on incorrect amine

ALLOW

$$H_3C$$
 N
 H

DO NOT ALLOW if ring is connected to the N triple bond in the diazonium or if diazonium has a negative charge **ALLOW** one mark for correct displayed diazonium if alkyl group is not shown

ALLOW

ALLOW

for both marks

ALLOW

$$\begin{array}{c} \textbf{ALLOW} \\ \text{H}_3C \\ \hline \end{array}$$

for one mark

HNO₂ + HCl and temp < 10 °C **OR** NaNO₂ + HCl and temp < 10 °C \checkmark alkaline **AND** phenol (if temperature stated must be below 10 °C) \checkmark **ALLOW** NaOH **OR** KOH & C₆H₅OH **OR** phenoxide is

ALLOW NaOH **OR** KOH & C_6H_5OH **OR** phenoxide ion **OR** $C_6H_5O^-$

ALLOW reagents and conditions from the equations

[5]

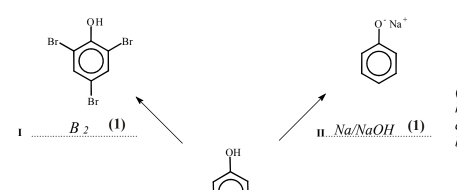
2

(ii) reducing agent (1)

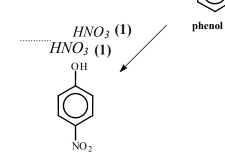
(c) moles $C_6H_5NO_2$ used = **0.0300** (mol) (**1**) theoretical yield of $C_6H_5NH_2$ = **2.79(3)** (g) (**1**) or ecf actual 72.1% yield = **2.014** (g) / (calculator value 2.013753) (**1**) or ecf to three sig figs = **2.01** (g) (**1**) or ecf

[9]

3. (i)



(do not allow a halogen carrier with the bromine)



(do not penalise use of a nitrating mixture)

(ii) dye / colouring / indicator (1)

1

5

3

(iii) phenylamine (1) $NaNO_2 / HNO_2$ (1) + HCl (1) < 10°C (1) add to alkaline phenol (1)

[9]

4. sodium nitrite + HCl / nitrous acid (1)

phenol/named example (added to the products from above) AW (1) alkaline conditions / OH^- (1)

example of an azo dye that could be formed from phenylamine,

6

[6]

- 5. (i) nitrous acid / HNO_2
 - (ii) $CH_3 \longrightarrow \stackrel{+}{N} = N \quad (C\Gamma) \quad (1)$
 - (iii) diazonium (ion /salt) (1)
 - (iv) to prevent decomposition / it reacting (diazonium ion) is unstable **AW**
 - (v) structure showing the amine coupled to the phenol or its salt e.g.

$$CH_3$$
 N
 N
 CH_3
 CH_3
 CH_3

-N=N-(1) rest of structure (joined by two nitrogens) (1)

[6]

1

1

6. methylation stage (can come anywhere)

 CH_3Cl / CH_3Br (1)

 $AlCl_3$ / FeBr₃ etc. (1)

equation – e.g. $C_6H_6 + CH_3Cl \rightarrow C_6H_5CH_3 + HCl$ (1)

intermediate name or unambiguous structure (1)

4 marks

intermediates and equations will vary if methylation is done after nitration or reduction

nitration stage

(conc) H₂SO₄ (1)

 $(conc) HNO_3 (1)$

equation – e.g.: $C_6H_5CH_3 + HNO_3 \rightarrow C_6H_4(CH_3)NO_2 + H_2O$ (1)

intermediate – name or unambiguous structure (1)

4 marks

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reduction stage
tin/iron (1)
HCl (1)
equation – e.g.: C_6H_4(CH_3)NO_2 + 6[H] \rightarrow C_6H_4(CH_3)NH_2 + 2H_2O
                 or with H<sup>+</sup> also on left to give C<sub>6</sub>H<sub>4</sub>(CH<sub>3</sub>)NH<sub>3</sub><sup>+</sup> (1)
3 marks
                     allow other suitable reducing agents:
Quality of Written Communication mark for a well organised
answer with the three stages clearly distinguished and sequenced (1)
1 mark
                                                                                                 12
       1<sup>st</sup> stage
(a)
       aromatic amine / named aromatic amine / structure (1)
       sodium nitrite / nitrous acid (1)
       HCl/H_2SO_4 (but not conc) /H^+ (1)
       at <10^{\circ}C (1)
       which forms a diazonium salt / ion (1)
                     if more than four are given, mark any wrong reagents,
                     conditions first
       2<sup>nd</sup> stage
       the product from the first stage mixed with the phenol AW (1)
       (in excess) hydroxide / alkali (1)
                                                                                                  7
                     allow correct formulae for the reagents
(b)
       (i)
                      ОН
                                                    (1)
                                                                                                   1
                     allow any benzene rings as well as N=N circled, as long as no
                     other groups are
              ...16... carbon and ......10..... hydrogen atoms
       (ii)
                (1)
                                         (1)
                                                                                                  2
```

7.

[12]

1

Plymstock School 5

Na / NaOH / OH⁻ etc (1)

(c)

(d)

OH

$$SO_3^- Na^+$$

(1)

 $allow \ I \ mark \ if \ they \ are \ both \ correct, \ but \ in \ the \ wrong \ boxes \ only \ penalise \ a \ slip \ with \ SO_3^- Na^+ \ once$

[13]

2

2

1

8. (a) (i) Diamino

two/2 amine groups (1)

1,4

their position on the ring / numbering of carbons around ring (or shown on a diagram) (1)

reduction / redox (1)

(b)

(i)

- (ii) tin and HCl (1) cone acid under reflux (1) 2 $or H_2 gas + Ni/Pd catalyst$

(iii)
$$O_2N \longrightarrow NO_2 + 12[H] \longrightarrow H_2N \longrightarrow NH_2 + 4H_2O$$

$$H_2O \text{ as product (1)}$$
and the equation balanced (1)

- (c) (i) accepts H⁺ using the lone pair (on N) (1) which is donated/forms a (dative) covalent bond (1) 2 either mark can be obtained with a good diagram
 - (ii)

 Cl H₃N[±] Cl Correct structure with charges shown (1)(1)

 one mark for either: just one neutralised,

 both neutralised, but without Cl,

 both neutralised, but no charges shown

 2

(iii) hexane-1,6-diamine is a stronger base because:

electrons move towards the N (due to the inductive effect) (in hexane-1,6-diamine) (1)

the lone $\underline{\text{pair}}$ from N is (partially) delocalised around the ring (in diaminobenzene) (1)

so the electron pair is more easily donated $\!\!/$

H⁺ more easily accepted (in hexane-1,6 diamine) ora (1)

[14]

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