²**Synthesis of Benzil by Air Oxidation of Benzoin** ³**and M(Salen) Catalyst**

9 **ABSTRACT**

Bis salicylaldehyde ethylenediamine Schiff base(Salen) and its complexes with three metal ions(Co2+, Ni2+, Zn2+) were prepared, and characterized by infrared spectroscopy(IR). Using air as oxygen source, the optimum reaction conditions for the catalytic oxidation of 0.05 mol benzoin by Co(Salen) were obtained by orthogonal test as follows: base KOH 2 g, catalyst 1.5 g, N, N-dimethylformamide(DMF) as solvent, reaction temperature 40 °C, reaction time 1 h. Under these conditions, the catalytic performances of different metal complexes were investigated. The catalytic activity of Co(Salen) was the best one, the yield of benzil was up to 93.6%, the number of Ni(Salen) and Zn(Salen) was 86.3% and 82.1%, respectively. The reused catalytic performance of M(Salen) complex was also studied. The catalytic activity of Co(Salen), Ni(Salen) and Zn(Salen) was stable after 4 times recycle, the yield of benzil was 71.4%, 63.3% and 57.4%, respectively, and it was easy for catalyst recycling. The oxidation product was certainly benzil with high purity according to the characterization results of melting point(MP), IR, high performance liquid chromatography(HPLC) and 1H nuclear magnetic resonance(1H NMR). Compared with the common synthetic method of benzil, this one has the advantages of friendly environment, low cost and easy operation. It is a simple and green way to synthesize benzoyl efficiently.

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12 *Keywords: Benzil; benzoin; M(Salen) catalyst; air oxidation; orthogonal experiment.*

14 **1. INTRODUCTION**

15 16 Benzil is an important organic intermediate and organic chemical raw material. It is 17 synthesized by oxidation of benzoin and widely used in food, medicine, pesticide and other industries.

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There are two methods for synthesizing benzil by oxidation of benzoin: one is non-catalytic 21 oxidation. The benzil is prepared by oxidizing benzoin with an incrganic compound or an $\frac{21}{2}$ oxidation. The benzil is prepared by oxidizing benzoin with an inorganic compound, or an 22 organometallic compound or a polymer, but there are problems such as large requirement. $\frac{22}{2}$ organometallic compound, or a polymer, but there are problems such as large requirement 23 of oxident, intense reaction, large amount of by products, serious pollution and so on. For $\frac{23}{2}$ of oxidant, intense reaction, large amount of by-products, serious pollution and so on. For 24 axample, Irene Dip et al. [1] reported that benzil was prepared from benzoin with ethyl example, Irene Dip et al [1] reported that benzil was prepared from benzoin with ethyl
25 acetate used as solvent 40% trichloro isocyanuric acid(TCCA) as oxidant the reaction time acetate used as solvent, 40% trichloro isocyanuric acid(TCCA) as oxidant, the reaction time 26 was as long as 24 h at room temperature, and the consumption of oxidant was greater was as long as 24 h at room temperature, and the consumption of oxidant was greater.

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The other is catalytic oxidation of benzil by using high efficiency catalyst and molecular 29 exigen or air as exident. This mothed has the educations of low environmental pollution $\frac{29}{20}$ oxygen or air as oxidant. This method has the advantages of low environmental pollution $\frac{29}{20}$ and conforms to the now concept of green chamista, and clean production. Stower A $\frac{30}{31}$ and conforms to the new concept of green chemistry and clean production. Steven A $\frac{31}{31}$ Tymonko at al. [2] reported the establise ovidation of benzein to benzil by using air as ovidant $\frac{31}{2}$ Tymonko et al [2] reported the catalytic oxidation of benzoin to benzil by using air as oxidant, 32 $\frac{32}{10}$ acetic acid aqueous solution as solvent, 40 mol% Bi(NO₃)₂ and 4 mol% Cu(Ac)₂ as catalyst, 33 the reaction could be carried out for only 3 b, the reaction conditions were mild and the the reaction could be carried out for only 3 h, the reaction conditions were mild and the 34 proporation of catalyst was simple, but it raised the set due to large Bi(NO), eatplyst ³⁴ preparation of catalyst was simple, but it raised the cost due to large Bi($NO₃$)₂ catalyst 35 decade. Tabira Shamim at al [3] reported that air as exident to use as solvent. Bd/SiO as dosage. Tahira Shamim et al [3] reported that air as oxidant, toluene as solvent, $Pd/SiO₂$ as

 $\frac{36}{37}$ a catalyst to oxidize benzoin to benzil at 100 °C, the reaction only needs 0.75 h, the reaction 37 time was short but the eatalyst was expensive and the reaction set was high which is not $\frac{37}{3}$ time was short, but the catalyst was expensive and the reaction cost was high, which is not 38 conductive to the industrial production conducive to the industrial production. 39

40 Therefore, it is of great significance to develop a kind of catalyst with low environmental 41 pollution, low cost and high reaction efficiency, which can selectively catalyze benzoin to 42 benzil under mild conditions. Schiff bases are a very important class of nitrogen-containing 43 ligands that work well with metal ions. Compared with mono Schiff base, the bis Schiff base 44 contains more coordination sites and forms more stable complexes with metals, such as bis
45 salicylaldehyde ethylenediamine(Salen) as ligand, which is often used to study the oxygensalicylaldehyde ethylenediamine(Salen) as ligand, which is often used to study the oxygen-46 carrying capacity and photothermal properties of complexes [4, 5].

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48 In this paper, from the perspective of green chemistry, M(Salen), the bis salicylaldehyde 49 ethylenediamine-metal complex was used as a catalyst to study the green synthesis of 50 benzil from benzoin. Using air as the oxidant, so the input cost is reduced, and the reaction 51 conditions are mild, the reaction process is green and environmentally friendly.

53 **2. EXPERIMENTAL DETAILS**

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55 **2.1 Reagents and instruments**

56 57 Reagents: benzoin(AR), salicylaldehyde(CP), ethylenediamine(AR), cobalt acetate(AR), zinc 58 acetate(AR), nickel acetate(AR), potassium hydroxide(AR), anhydrous ethanol(AR), N, N-59 dimethylformamide(DMF, AR), dichloromethane(AR), anhydrous magnesium sulfate(AR), 60 benzil(purity 98%).

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Instruments: Thermo Fisher NICOLET ISIO Fourier infrared spectrometer, Hitachi LC-2130 63 liquid chromatograph, WRS-1A melting point instrument, 78HW-1 digital thermostat 64 magnetic stirrer, ZX98-1 rotary evaporator, Bruker Advance III HD 500MHz nuclear magnetic 65 resonance instrument.

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67 **2.2 Synthesis of M(Salen) catalyst**

- 68
- 69 The synthesis of the M(Salen) complex catalyst is shown in the following scheme 1: 70

bis salicylaldehyde ethylenediamine-Mcomplex[M(Salen)]

73 **Scheme 1 Synthesis of M(Salen) catalyst**

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75 The process is as follows: 0.10 mol salicylaldehyde and 60 mL anhydrous ethanol were 76 added to the three-necked flask, 0.05 mol ethylenediamine was further added under stirring, 77 and the mixture was heated and refluxed for 1 h. Then, a 35 mL aqueous solution containing 78 0.05 mol CoAc₂ was dropped into the three-necked flask, and the solution was stirred at 75 79 °C for 50 min. After cooling, suction filtration was carried out to obtain a dark red solid, which 80 was washed and dried to give bis salicylaldehyde ethylenediamine-cobalt 81 complex[Co(Salen)][6].

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83 Using the same method as above, Ni(Salen)(earth yellow solid), Zn(Salen)(light yellow solid) 84 were prepared by the combination of NiAc₂, ZnAc₂ with bis salicylaldehyde ethylenediamine, 85 respectively. respectively. 86

87 **2.3 Catalytic oxidation of benzoin** 88

89 The catalytic oxidation of benzoin by M(Salen) is shown in the following scheme 2: 90

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Scheme 2 Catalytic oxidation of benzoin

95 A typical reaction is: 0.05 mol benzoin and 60 mL DMF were added to a three-necked flask 96 equipped with a stirrer, reflux condenser and air duct. After dissolving, 1.5 g Co(Salen) 97 catalyst and 2 g KOH were added, air was pumped in to oxidize benzoin. The solution was 98 heated to 40 °C in water bath, and the reaction process was tracked by thin layer 99 chromatography (TLC). After the completion of the oxidation reaction, the mixture was 100 cooled to room temperature, and the pH of the reaction solution was adjusted to 3-4. The 101 mixture was poured into 150 mL water to precipitate a solid, which was filtered with suction 102 and washed with water, then the yellow needle crystal benzil was obtained.

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104 **2.4 Recycling of M(Salen) catalyst**

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106 The filtrate after the catalytic oxidation reaction of benzoin was extracted with CH_2Cl_2 in 107 portions, and the extract was dried with anhydrous magnesium sulfate. The CH₂Cl₂ was 107 portions, and the extract was dried with anhydrous magnesium sulfate. The CH_2Cl_2 was 108 recovered by rotary evaporator, and the residue was a DMF solution containing M(Salen) recovered by rotary evaporator, and the residue was a DMF solution containing M(Salen) 109 catalyst. A small amount of DMF was added, fresh benzoin and KOH were added to carry 110 out the next batch of catalytic oxidation reaction, so that the M(Salen) catalyst was reused.

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112 **3. RESULTS AND DISCUSSION** 113

114 **3.1 Infrared spectrum analysis of M(Salen) catalyst**

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116 The infrared spectrum of M(Salen) is shown in Fig. 1. The three catalysts all have absorption 117 peaks caused by the vibration of the benzene ring skeleton at 1600-1630 cm⁻¹, and by the 118 stretching vibration of a bidentate chelating bond(two oxygen atoms coordinated with the 119 surface M cation simultaneously) at 1340-1350 cm⁻¹. The absorption peak at 1150-1200 cm⁻¹ 120 and 1050-1090 cm^{-1} are caused by the C-N and C-O stretching vibration of the Salen ligand, the peak at 740-750 cm⁻¹ corresponds to the out-of-plane bending vibration of C-H of the 122 benzene ring. The above results indicated that three kinds of M(Salen) catalysts had been 123 successfully prepared.

128 **3.2 Orthogonal experiment to explore the optimal reaction conditions for** 129 **catalytic oxidation of benzoin**

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131 Co(Salen) used as catalyst, a series orthogonal experiments with 5-factor and 4-level were
132 designed to study the effect on benzoin catalytic oxidation to benzil. Reaction 132 designed to study the effect on benzoin catalytic oxidation to benzil. Reaction 133 temperature(A), reaction time(B), base KOH amount(C), catalyst amount(D) and solvent 134 type(E) were five factors, each factor had four levels. The factors and corresponding levels 135 of $L_{16}(4^5)$ orthogonal experiment is listed in Table 1. of $L_{16}(4^5)$ orthogonal experiment is listed in Table 1.

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Table 1 The factors and levels of this $L_{16}(4^5)$ **orthogonal experiment**

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139 According to the design principle of orthogonal experiment, we designed the factor-level 140 scheme of Table 2. The benzoin catalytic oxidation experiments were carried out according 141 to the reaction conditions arranged in Table 2, and the yield of benzil was used as an 142 experimental index.

Table 2 Factor-level scheme and experimental results of L₁₆(4⁵) orthogonal experiment

No.	Reaction temperature (°C)	Reaction time (h)	KOH amount (g)	Co(Salen) amount (g)	Solvent	Benzil yield (%)
	20	0.5		0.5	100% EtOH	62.0
っ	20				80% EtOH	87.9
3	20	1.5		1.5	60% EtOH	75.7
	20				DMF	77 1
5	40	0.5		1.5	DMF	89.7

146 From the yield data, we can obtain the data analysis in Table 3. From the mean value of 147 each level labeled as K_i , the optimum levels of each factor can be judged. The maximum K_i 148 value of each factor was: 77.6(K_2 of A), 79.4(K_2 of B), 81.1(K_2 of C), 79.1(K_3 of D) and 149 80.1(K_4 of E), respectively. So the optimal reaction conditions of the experiment obtained $80.1(K₄$ of E), respectively. So the optimal reaction conditions of the experiment obtained 150 were as follows: the reaction temperature was 40 °C(the 2^{nd} level), the reaction time was 1 151 h(the 2nd level), the amount of KOH was 2 g(the 2nd level), the amount of Co(Salen) catalyst 152 was 1.5 g(the $3rd$ level) and the solvent was DMF(the 4th level).

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154 The data of range labeled as R was 3(A), 5.4(B), 9.7(C), 6(D) and 9.1(E), respectively. This 155 indicated that the amount of base(factor C) had the greatest influence on the catalytic 156 oxidation of benzoin, followed by the solvent(factor E), then the amount of catalyst(factor D) 157 and the reaction time(factor B), the reaction temperature(factor A) had the least influence on 158 it. That is to say, the importance of each factor effect on benzoin catalytic oxidation to benzil 159 was base amount > solvent type > catalyst amount > reaction time > reaction temperature.

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161 **Table 3 Analysis of orthogonal experimental results**

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163 **3.3 Catalytic performance of different M(Salen)**

165 According to the optimal reaction conditions obtained above, we used Co(Salen), Ni(Salen) 166 and Zn(Salen) as catalysts to carry out the oxidation reaction of benzoin. The yield of the 167 obtained benzil is shown in Table 4. It can be seen that the catalytic effect of Co(Salen) was 168 the best one, the yield of benzil was 93.6%. The yield of product obtained by catalytic 169 oxidation using Ni(Salen) was in close proximity to that of Zn(Salen), and the value was 170 86.3% and 82.1%, respectively.

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172 **Table 4 Catalytic performances of different Catalysts**

vst Catal	Co (Salen)	ılen)	len) フロハ
Benzil yield(%)	93.6	າຂາ	
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173 **Reaction conditions: benzoin 0.05 mol, KOH 2 g, M(Salen) catalyst 1.5 g, DMF 60 mL, reaction* temperature 40 °C, reaction time 1 h.

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176 **3.4 Recycling performance of M(Salen) catalyst**

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178 According to the above optimal reaction conditions, we used Co(Salen), Ni(Salen) and 179 Zn(Salen) as catalysts to carry out the recovery and reuse performance studies. The yield of 180 the obtained benzil is shown in Table 5.

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182 It can be seen from Table 5 that with the increase of the number of the reuse of the catalyst, 183 the catalytic activity gradually decreased, but when the catalyst was used for the fourth time, 184 the benzil yield on Co(Salen), Ni(Salen) and Zn(Salen) catalysts was still maintained at 185 71.4%, 63.3% and 57.4%, respectively. The better recovery and reuse performance of 186 M(Salen) can reduce the economic cost of the catalyst.

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189 **Reaction conditions: benzoin 0.05 mol, KOH 2 g, M(Salen) catalyst 1.5 g, DMF 60 mL, reaction* temperature 40 °C, reaction time 1 h.

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192 **3.5 Characterization of the benzil product**

193 194 The obtained product benzil was first determined by a melting point(MP) analyzer, and the 195 melting range was 94.2-94.7 °C, which was consistent with the theoretical melting point of 196 benzil.

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198 The product was further characterized by infrared spectroscopy(IR), high performance liquid 199 chromatography(HPLC) and ¹H nuclear magnetic resonance(¹H NMR).

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201 Fig. 2 is the infrared spectrum of benzil, which is similar to the standard spectrum in the 202 infrared spectrometer's own database. The absorption peak at 3063 cm⁻¹ corresponds to the 203 C-H stretching vibration of the methylene group, the peak at 1659 cm⁻¹ corresponds to the 204 C=O stretching vibration of carbonyl group. The carbonyl group is conjugated with the 205 benzene ring, so the absorption shifts to a low frequency(the normal absorption frequency of 206 C=O is at 1740-1700 cm^{-1}). 1593 cm⁻¹, the absorption peak at this position corresponds to the 207 vibration of the benzene ring skeleton, the strong peak at 1211 cm^{-1} corresponds to the 208 stretching vibration of C-C, and the peak at 718 cm^{-1} corresponds to the out-of-plane 209 bending vibration of C-H on the benzene ring.

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214 The qualitative analysis and purity of the obtained oxidation product benzil was carried out 215 by HPLC using an Eclipse Plus C18 column, 70% methanol as the mobile phase, flow rate by HPLC using an Eclipse Plus C18 column, 70% methanol as the mobile phase, flow rate 216 1.0 mL/min, column temperature 35 °C, detection wavelength 259 nm, and injection quantity
217 5 uL. The standard curve shown in Fig. 3 was obtained by taking the concentration of benzil 217 5 μL. The standard curve shown in Fig. 3 was obtained by taking the concentration of benzil
218 standard solution series $(x10^{-6})$ as 40, 80, 200, 400, 600, 800, 1000 mol/L. The regressive standard solution series $(*10^{-6})$ as 40, 80, 200, 400, 600, 800, 1000 mol/L. The regressive 219 equation and correlation coefficient were $Y=266016+5991X$ and $R^2=0.9916$, respectively, 220 indicating the standard curve in the range of 4×10^{-5} -1 $\times 10^{-3}$ mol/L had good linear 221 relationship. 222

223 Then, the reaction product benzil 0.0105 g was weighed, and the solution was adjusted to 224 99.89 \times 10⁻⁶ mol/L after dilution. The solution was sampled, the qualitative and the 225 quantitative analyses were carried out based on the retention time and peak area
226 respectively. The retention time of the product peak was found to be consistent with the respectively. The retention time of the product peak was found to be consistent with the 227 retention time of the standard solution peak $(t = 6.5 \text{ min})$, confirming that the product was 228 indeed benzil. By substituting the peak area $(Y=857272)$ into the standard curve, the actual 228 indeed benzil. By substituting the peak area (Y=857272) into the standard curve, the actual concentration of the oxidized product benzil was 98.69×10^{-6} mol/L, so the purity of the benzil concentration of the oxidized product benzil was 98.69×10^{-6} mol/L, so the purity of the benzil 230 product was 98.69×10^{-6} /99.89 \times 10⁻⁶ =98.80%.

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233 **Fig.3. LC standard curve of benzil**

234 235 The obtained oxidation product benzil was also characterized by 1H NMR, and the result is 236 shown in Fig. 4. The peak with a chemical shift of 7.99-7.50 ppm corresponds to hydrogen 237 on the benzene ring. According to the peak area data, the ratio of three kinds of hydrogen is 238 2.10: 1.00: 2.19, which is close to 2: 1: 2 in accordance with the molecular formula of benzil, 239 further confirming that the synthesis product was indeed benzil. 240

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244 **4. CONCLUSION**

246 We prepared M(Salen) complexes and used it for the air oxidation synthesis of benzil from 247 benzoin. The experimental results showed that the M(Salen) complex has a good catalytic 248 effect.

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250 Orthogonal test method was carried out to investigate the optimum reaction conditions of 251 0.05 mol benzoin catalyzed by Co(Salen). The optimal reaction conditions were as follows: 252 base KOH amount 2 g, catalyst amount 1.5 g, solvent DMF, reaction temperature 40 $^{\circ}$ C, 253 reaction time 1 h. The yield of benzil was as high as 93.6%, and the yield of the product 254 catalyzed by Ni(Salen) and Zn(Salen) was also up to 86.3% and 82.1%, respectively.

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256 The use of air as an oxidant reduces the input cost of reaction, and the reaction device is 257 simple, the operation is also convenient. Meanwhile, the M(Salen) complex is easy to 258 prepare, with the advantages as less dosage, high catalytic efficiency and convenient post-259 treatment. The recovered catalyst can be reused, reducing the production cost and meeting 260 the requirements of green chemistry. After four cycles of use, the yield of benzil on the

261 Co(Salen), Ni(Salen) and Zn(Salen) catalyst still reached 71.4%, 63.3% and 57.4%, respectively.

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264 The oxidized product benzil was characterized by MP, IR, HPLC and 1 H NMR, all results of 265 which proved the product was certainly high purity benzil. It was confirmed that M(Salen) 266 complex catalyze the oxidation of benzoin to benzil was green and feasible.

267 268

269 **COMPETING INTERESTS** 270

271 Authors have declared that no competing interests exist. 272

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