

## LAMPIRAN A

### CONTOH PERHITUNGAN PERSENTASE HASIL SINTESIS

#### I. Perhitungan berat teoritis

a. Asam Antranilat (BM : 137,14 g/mol)

Penimbangan : 13,7 gram

$$\text{mol asam antranilat} : \frac{13,7}{137,14} = 0,1 \text{ mol}$$

b. *p*-klorobenzoil klorida (BM : 175,02 g/mol, berat jenis : 1,377 g/cm<sup>3</sup>)

Volume : 25,42 ml

$$\text{mol } p\text{-klorobenzoil klorida} : \frac{25,42 \times 1,377}{175,02} = 0,2 \text{ mol}$$

c. Hidrazin Hidrat (BM : 50,05 g/mol, berat jenis : 1,03 g/cm<sup>3</sup>)

Volume : 2,43 ml

$$\text{mol hidrazin hidrat} : \frac{2,43 \times 1,03}{50,05} = 0,05 \text{ mol}$$

d. Benzaldehida (BM : 106,12 g/mol, berat jenis : 1,05 g/cm<sup>3</sup>)

Volume : 2,03 ml

$$\text{mol benzaldehida} : \frac{2,03 \times 1,05}{106,12} = 0,02 \text{ mol}$$

## II. Perhitungan persentase hasil sintesis berdasarkan mmol teoritis

Persentase hasil 2-(*p*-klorofenil)-4*H*-3,1-benzoksazin-4-on :



awal	0,01 mol		0,02 mol		
reaksi	0,01 mol	+	0,01 mol	-	0,01 mol
sisa	0		0,01 mol		0,01 mol

$$\text{BM teoritis} = 359$$

$$\text{Massa teoritis} = 0,01 \text{ mol} \times 359 = 3,59 \text{ gram}$$

$$\text{Massa praktis} = 2,66 \text{ gram}$$

$$\% \text{ hasil} = \frac{2,66}{3,59} \times 100\% = 74 \%$$

**LAMPIRAN B**  
**PERHITUNGAN UJI T STATISTIK**

**Pengaruh Penambahan 2-klorobenzaldehid Dibanding Benzaldehid :**

---

t-Test: Paired Two Sample for Means

	<i>Variable 1</i>	<i>Variable 2</i>
Mean	72.66666667	70
Variance	2.333333333	1
Observations	3	3
Pearson Correlation	0.654653671	
Hypothesized Mean Difference	0	
df	2	
t Stat	4	
P(T<=t) one-tail	0.028595479	
t Critical one-tail	2.91998558	
P(T<=t) two-tail	0.057190958	
t Critical two-tail	4.30265273	

---

## Pengaruh Penambahan 2,4-diklorobenzaldehid Dibanding Benzaldehid

t-Test: Paired Two Sample for Means

	Variable 1	Variable 2
Mean	72.66666667	66
Variance	2.333333333	1
Observations	3	3
Pearson Correlation	-0.327326835	
Hypothesized Mean Difference	0	
df	2	
t Stat	5.547001962	
P(T<=t) one-tail	0.015498417	
t Critical one-tail	2.91998558	
P(T<=t) two-tail	0.030996834	
t Critical two-tail	4.30265273	

**LAMPIRAN C**  
**HARGA Rf OPTIMASI SENYAWA**

- a. Harga Rf optimasi sintesis senyawa 2-(p-klorofenil)-3,1-benzoksazin-4(3H)-on

Senyawa	Harga Rf
Asam antranilat	0,48
<i>p</i> -klorobenzoil klorida	0,30
Sampel menit ke-30	0,48 & 0,30
Sampel menit ke-45	0,48 & 0,30
Sampel menit ke-60	0,79 & 0,30
Sampel menit ke-75	0,79 & 0,30

- b. Harga Rf optimasi sintesis senyawa 3-amino-2-(p-klorofenil)kuinazolin-4(3H)-on

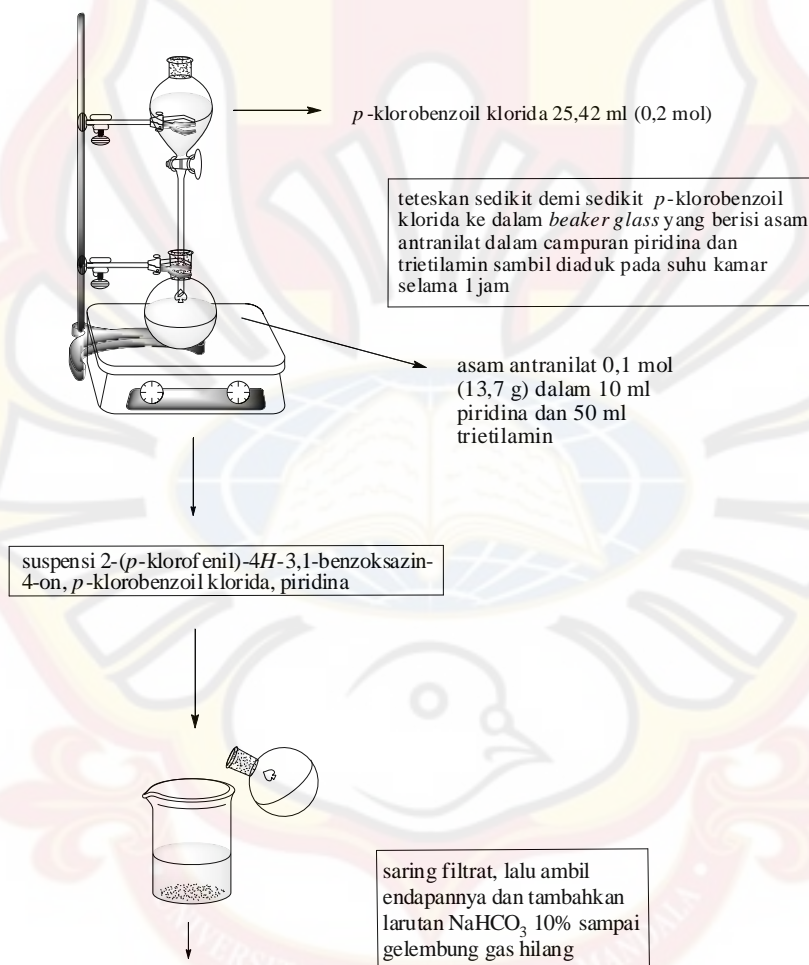
Senyawa	Harga Rf
2-( <i>p</i> -klorofenil)-3,1-benzoksazin-4(3H)-on	0,71
Sampel jam ke-1	0,70
Sampel jam ke-2	0,71
Sampel jam ke-3	0,71
Sampel jam ke-4	0,70
Sampel jam ke 5	0,38

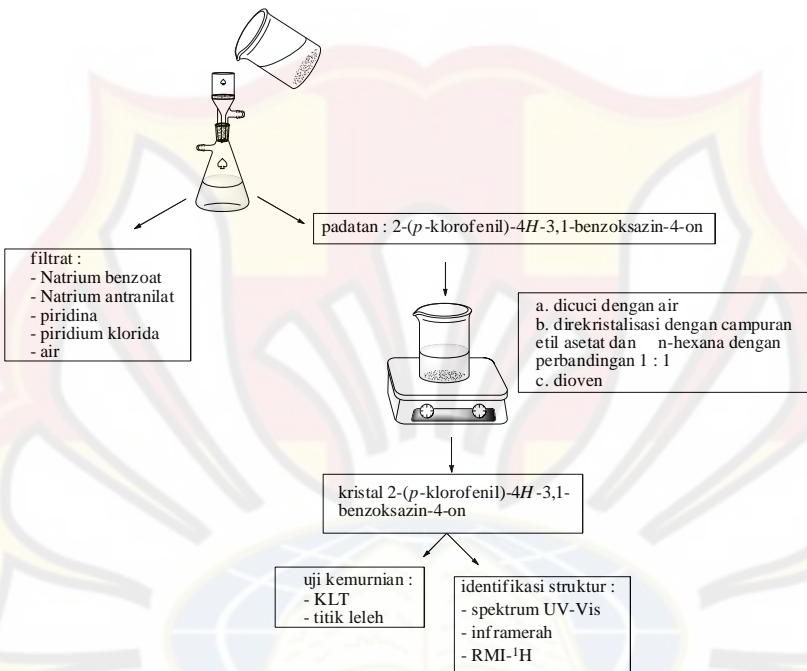
- c. Harga Rf optimasi sintesis senyawa 3-benzilidenamino-2-(p-klorofenil)kuinazolin-4(3H)-on

Senyawa	Harga Rf
3-amino-2-( <i>p</i> -klorofenil)kuinazolin-4(3 <i>H</i> )-on	0,71
Benzaldehida	0,74
Sampel menit ke-30	0,71 & 0,74
Sampel menit ke-45	0,70 & 0,74
Sampel menit ke-60	0,71 & 0,74
Sampel menit ke-75	0,81 & 0,74

## LAMPIRAN D

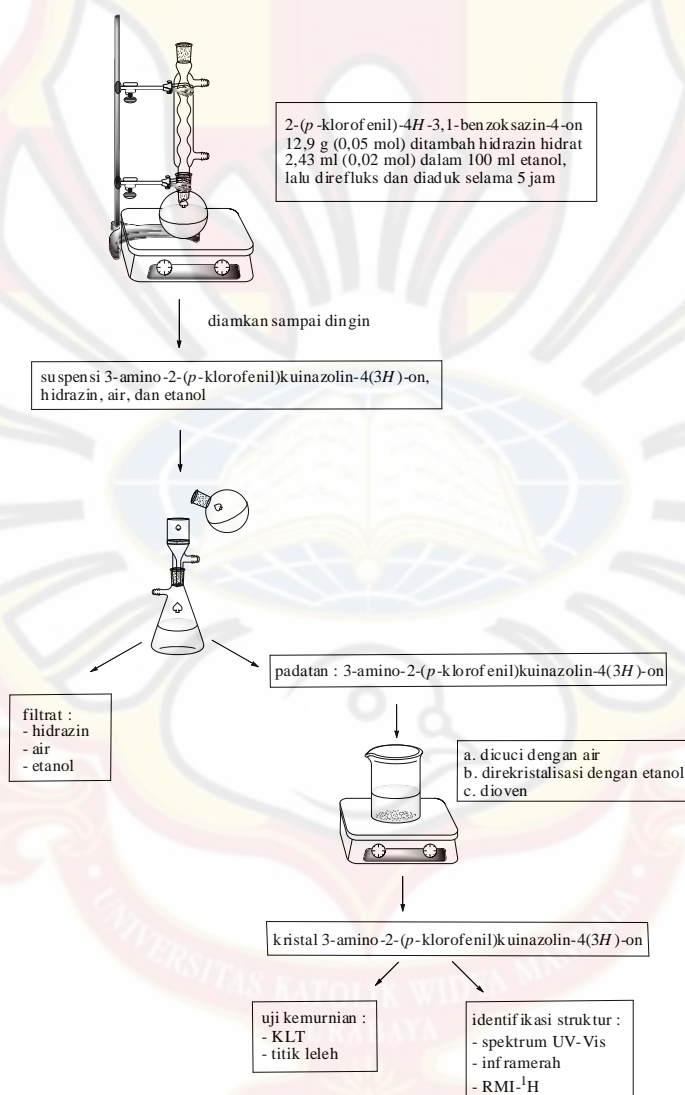
### SKEMA KERJA SINTESIS 2-(*p*-KLOROFENIL)-4H-3,1-BENZOKSAZIN-4-ON





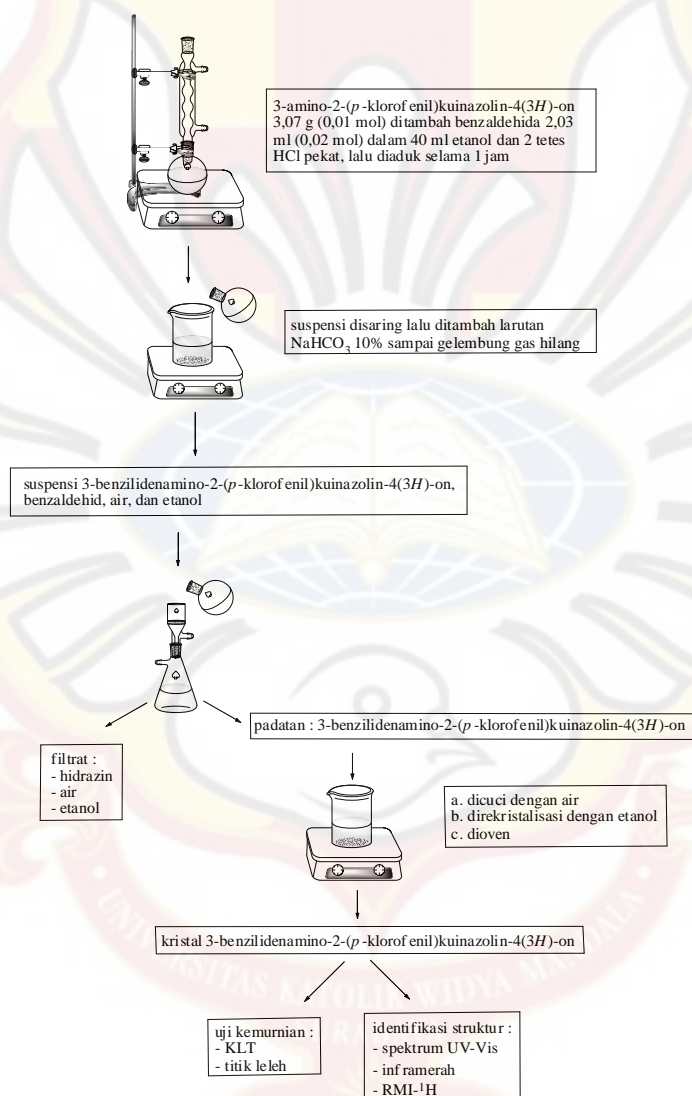
## LAMPIRAN E

### SKEMA KERJA SINTESIS 3-AMINO-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3H)-ON



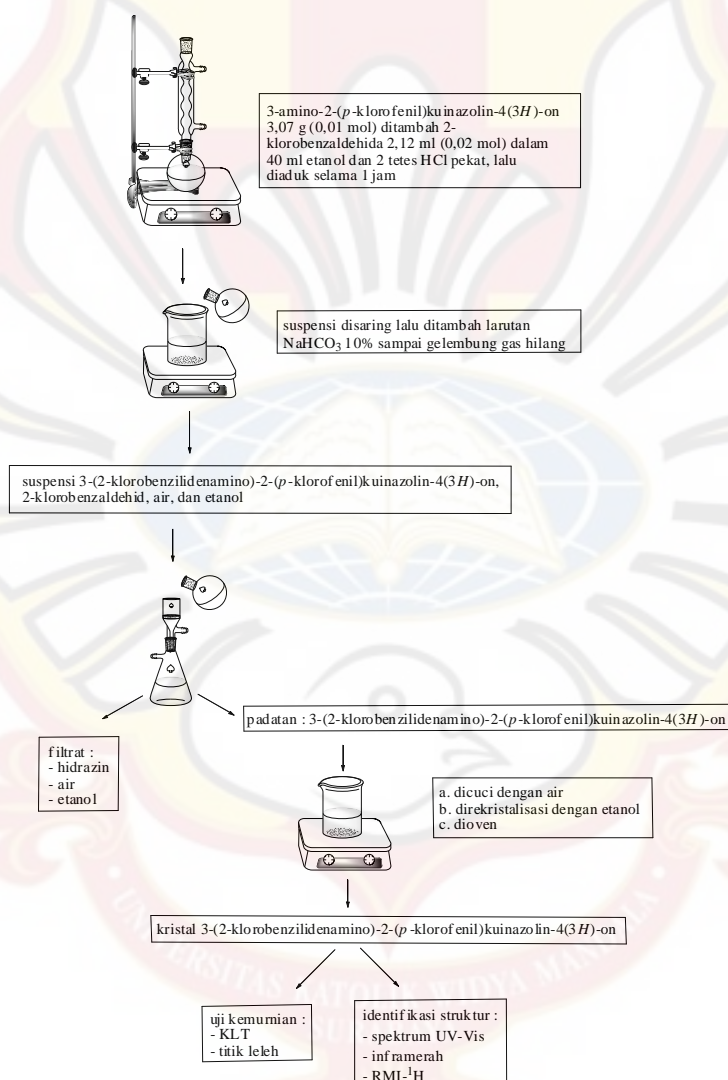
## LAMPIRAN F

### SKEMA KERJA SINTESIS 3-BENZILIDENAMINO-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3*H*)-ON



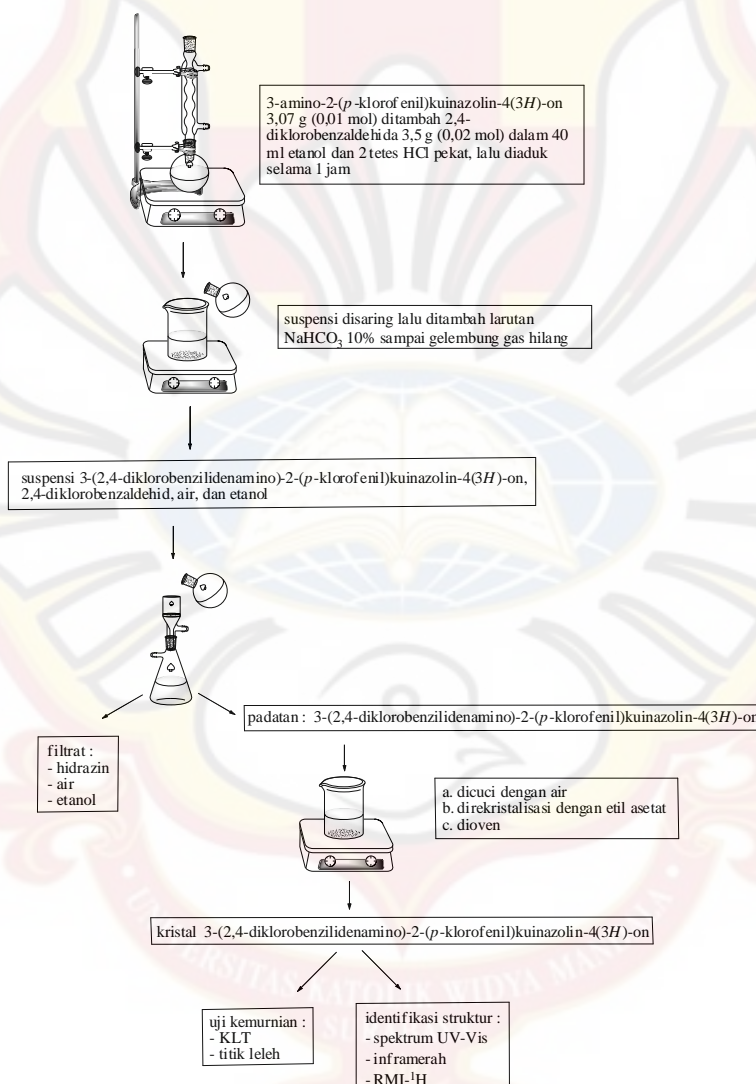
## LAMPIRAN G

### SKEMA KERJA SINTESIS 3-(2-KLOROBENZILIDENAMINO)-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3*H*)-ON



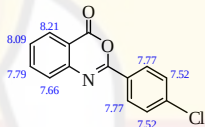
## LAMPIRAN H

### SKEMA KERJA SINTESIS 3-(2,4-DIKLOROBENZILIDENAMINO)-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3*H*)-ON

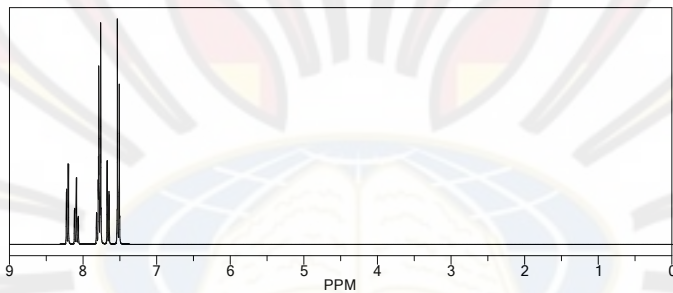


## LAMPIRAN I

### ESTIMASI RMI-<sup>1</sup>H 2-(*p*-KLOROFENIL)-4H-3,1-BENZOKSAZIN-4-ON



Estimation quality is indicated by color: **good**, **medium**, **rough**



Protocol of the H-1 NMR Prediction:

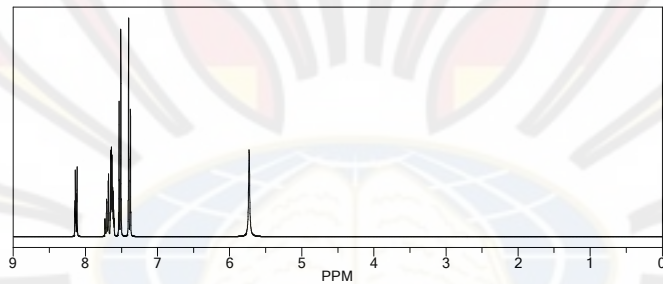
Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
CH 7.52	7.29	benzylidenimin	
	0.00	1-1C* <sup>6</sup> C* <sup>8</sup> C* <sup>3</sup> C* <sup>1</sup>	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.66	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.21	1-C(=O)O	
	0.19	general corrections	
CH 8.21	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.87	1-C(=O)O	
	0.08	general corrections	
CH 7.77	7.62	benzylidenimin	
	0.00	1-1C* <sup>6</sup> C* <sup>8</sup> C* <sup>3</sup> C* <sup>1</sup>	
	-0.06	1-Cl from 1-benzene	
	0.21	general corrections	
CH 7.52	7.29	benzylidenimin	
	0.00	1-1C* <sup>6</sup> C* <sup>8</sup> C* <sup>3</sup> C* <sup>1</sup>	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.77	7.62	benzylidenimin	
	0.00	1-1C* <sup>6</sup> C* <sup>8</sup> C* <sup>3</sup> C* <sup>1</sup>	
	-0.06	1-Cl from 1-benzene	
	0.21	general corrections	
CH 7.79	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.34	1-C(=O)O	
	0.19	general corrections	
CH 8.09	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.21	1-C(=O)O	
	0.62	general corrections	

## LAMPIRAN J

### ESTIMASI RMI-<sup>1</sup>H 3-AMINO-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3H)-ON



Estimation quality is indicated by color: **good**, **medium**, **rough**

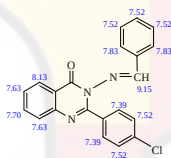


Protocol of the H-1 NMR Prediction:

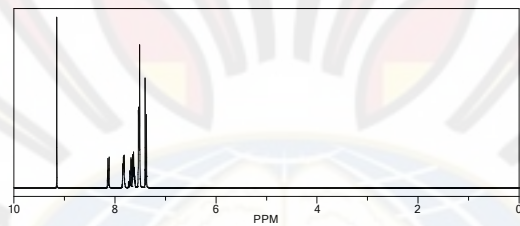
Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
NH2 5.73	2.00	amine	
	3.73	general corrections	
CH 7.52	7.29	benzylideminin	
	0.00	1-1*C=C*C=C*C=C*	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.63	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.18	1-C(=O)N	
	0.19	general corrections	
CH 8.13	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.69	1-C(=O)N	
	0.18	general corrections	
CH 7.39	7.62	benzylideminin	
	0.00	1-1*C=C*C=C*C=C*	
	-0.06	1-Cl from 1-benzene	
	-0.17	general corrections	
CH 7.52	7.29	benzylideminin	
	0.00	1-1*C=C*C=C*C=C*	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.39	7.62	benzylideminin	
	0.00	1-1*C=C*C=C*C=C*	
	-0.06	1-Cl from 1-benzene	
	-0.17	general corrections	
CH 7.70	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.25	1-C(=O)N	
	0.19	general corrections	
CH 7.63	7.26	1-benzene	
	?	1 unknown substituent(s)	
	0.18	1-C(=O)N	
	0.19	general corrections	

# LAMPIRAN K

## ESTIMASI RMI-<sup>1</sup>H 3-BENZILIDENAMINO-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3*H*)-ON



Estimation quality is indicated by color: good, medium, rough

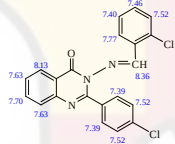


Protocol of the H-1 NMR Prediction:

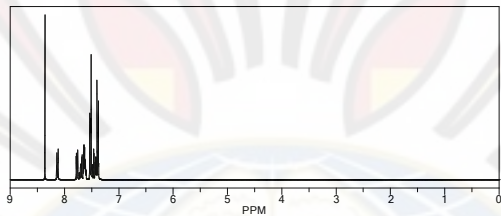
Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
CH 7.52	7.29	benzylideneimin	
	0.00	1-1C=C*C=C*C=C*	
	0.01	1-C(=O)N	
	0.22	general corrections	
CH 7.63	7.26	1-benzeno	
	?	1 unknown(s) substituent(s)	
	0.18	1-C(=O)O	
	0.19	general corrections	
CH 8.13	7.26	1-benzeno	
	?	1 unknown(s) substituent(s)	
	0.69	1-C(=O)N	
	0.18	general corrections	
CH 7.39	7.62	benzylideneimin	
	0.00	1-1C=C*C=C*C=C*	
	-0.01	1-CI from 1-benzeno	
	-0.17	general corrections	
CH 7.83	7.62	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	0.21	general corrections	
CH 7.52	7.29	benzylideneimin	
	0.00	1-1C=C*C=C*C=C*	
	0.01	1-CI from 1-benzeno	
	0.22	general corrections	
CH 7.39	7.62	benzylideneimin	
	0.00	1-1C=C*C=C*C=C*	
	-0.01	1-CI from 1-benzeno	
	-0.17	general corrections	
CH 7.83	7.62	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	0.21	general corrections	
CH 7.52	7.29	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	0.23	general corrections	
CH 7.70	7.26	1-benzeno	
	?	1 unknown(s) substituent(s)	
	0.25	1-C(=O)N	
	0.19	general corrections	
CH 7.63	7.26	1-benzeno	
	?	1 unknown(s) substituent(s)	
	0.18	1-C(=O)O	
	0.19	general corrections	
CH 7.52	7.29	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	0.23	general corrections	
CH 7.52	7.29	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	0.23	general corrections	
CH 9.15	8.11	benzylideneimin	
	?	1 unknown(s) substituent(s)	
	1.04	general corrections	

## LAMPIRAN L

### ESTIMASI RMI-<sup>1</sup>H 3-(2-KLOROBENZILIDENAMINO)-2-(p-KLOROFENIL)KUINAZOLIN-4(3H)-ON



Estimation quality is indicated by color: good, medium, rough

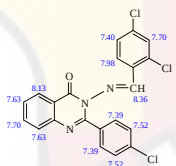


Protocol of the H-I NMR Prediction:

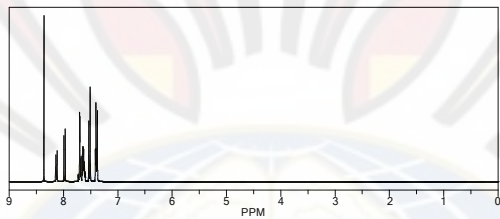
Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
CH 7.52	7.29	benzildiamin	
	0.00	1-C(=C\*)C\*C=C\*C=C\*1	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.63	7.26	1-benzene	
	? 0.18	1 unknown substituent(s)	
	0.19	general corrections	
CH 8.13	7.26	1-benzene	
	? 0.69	1 unknown substituent(s)	
	0.18	general corrections	
CH 7.39	7.62	benzildiamin	
	0.00	1-C(=C\*)C\*C=C\*C=C\*1	
	-0.06	1-Cl from 1-benzene	
	-0.17	general corrections	
CH 7.52	7.29	benzildiamin	
	0.00	1-C(=C\*)C\*C=C\*C=C\*1	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.52	7.29	benzildiamin	
	? 0.01	1 unknown substituent(s)	
	0.01	1-Cl from 1-benzene	
	0.22	general corrections	
CH 7.39	7.62	benzildiamin	
	0.00	1-C(=C\*)C\*C=C\*C=C\*1	
	-0.06	1-Cl from 1-benzene	
	-0.17	general corrections	
CH 7.77	7.68	benzildiamin	
	? 0.06	1 unknown substituent(s)	
	-0.06	1-Cl from 1-benzene	
	0.21	general corrections	
CH 7.46	7.29	benzildiamin	
	? 0.01	1 unknown substituent(s)	
	-0.06	1-Cl from 1-benzene	
	0.23	general corrections	
CH 7.40	7.29	benzildiamin	
	? 0.12	1 unknown substituent(s)	
	-0.12	1-Cl from 1-benzene	
	0.23	general corrections	
CH 7.70	7.26	1-benzene	
	? 0.25	1 unknown substituent(s)	
	0.19	1-C(=O)N	
	0.19	general corrections	
CH 7.63	7.26	1-benzene	
	? 0.18	1 unknown substituent(s)	
	0.19	1-C(=O)N	
	0.19	general corrections	
CH 8.36	8.11	benzildiamin	
	? 0.25	1 unknown substituent(s)	
	0.25	general corrections	

## LAMPIRAN M

### ESTIMASI RMI-<sup>1</sup>H 3-(2,4-DIKLOROBENZILIDENAMINO)-2-(*p*-KLOROFENIL)KUINAZOLIN-4(3H)-ON



Estimation quality is indicated by color: good, medium, rough



Protocol of the H-1 NMR Prediction:

Node	Shift	Base + Inc.	Comment (ppm rel to TMS)
CH 7.70	7.29	benzyldenimin	
	? 1 unknown substituent(s)		
	0.01 1-Cl from 1-benzene		
	0.01 1-Cl from 1-benzene		
	0.39 general corrections		
CH 7.52	7.29 benzylamin		
	0.00 1-[C=C=C=C=C=C]1		
	0.01 1-Cl from 1-benzene		
	0.22 general corrections		
CH 7.63	7.26 1-benzene		
	? 1 unknown substituent(s)		
	0.18 1-[C=O]N		
	0.19 general corrections		
CH 8.13	7.26 1-benzene		
	? 1 unknown substituent(s)		
	0.69 1-[C=O]N		
	0.18 general corrections		
CH 7.39	7.62 benzyldenimin		
	0.00 1-[C=C=C=C=C=C]1		
	-0.06 1-Cl from 1-benzene		
	-0.17 general corrections		
CH 7.52	7.29 benzyldenimin		
	0.00 1-[C=C=C=C=C=C]1		
	0.01 1-Cl from 1-benzene		
	0.22 general corrections		
CH 7.40	7.29 benzyldenimin		
	? 1 unknown substituent(s)		
	-0.12 1-Cl from 1-benzene		
	0.01 1-Cl from 1-benzene		
	0.22 general corrections		
CH 7.39	7.62 1-benzylamin		
	0.00 1-[C=C=C=C=C=C]1		
	-0.06 1-Cl from 1-benzene		
	-0.17 general corrections		
CH 7.98	7.62 benzyldenimin		
	? 1 unknown substituent(s)		
	-0.06 1-Cl from 1-benzene		
	-0.06 1-Cl from 1-benzene		
	0.48 general corrections		
CH 7.70	7.26 1-benzene		
	? 1 unknown substituent(s)		
	0.25 1-[C=O]N		
	0.19 general corrections		
CH 7.63	7.26 1-benzene		
	? 1 unknown substituent(s)		
	0.18 1-[C=O]N		
	0.19 general corrections		
CH 8.36	8.11 benzyldenimin		
	? 1 unknown substituent(s)		
	0.25 general corrections		