

## LAMPIRAN A

### CONTOH PERHITUNGAN PERSENTASE HASIL SINTESIS

#### I. Perhitungan berat teoritis

a. Ammonium tiosianat (BM : 76,12 g/mol)

Penimbangan : 1,14 gram

$$\text{mmol ammonium tiosianat} : \frac{1,14}{76,12} \times 1000 = 14,98 \text{ mmol} \cong 15 \text{ mmol}$$

b. 2-klorobenzoil klorida (BM : 175,01 g/mol, berat jenis : 1,378 g/cm<sup>3</sup>)

Volume : 1,3 ml

$$\text{mmol 2-klorobenzoil klorida} : \frac{1,3 \times 1,378}{175,01} \times 1000 = 10,2 \text{ mmol} \cong 10 \text{ mmol}$$

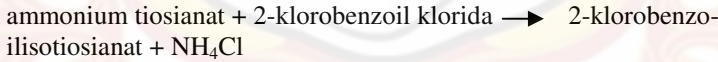
c. Amonia (BM : 35,05 g/mol, berat jenis : 0,91 g/cm<sup>3</sup>)

Volume : 1,15 ml

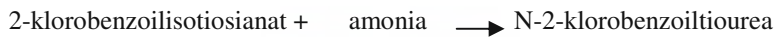
$$\text{mmol amonia} : \frac{1,15 \times 0,91}{35,05} \times 1000 = 30 \text{ mmol}$$

#### II. Perhitungan persentase hasil sintesis berdasarkan mmol teoritis

Persentase hasil N-2-klorobenzoiltiourea :



awal	15 mmol	10 mmol	0	0	
reaksi	10 mmol	10 mmol -	10 mmol	10 mmol	+
sisa	5 mmol	0	10 mmol	10 mmol	



awal	10 mmol	30 mmol	0	
reaksi	10 mmol	10 mmol -	10 mmol	+
sisa	0	20 mmol	10 mmol	

$$\text{BM teoritis} = 210,5$$

$$\text{Massa teoritis} = 10 \text{ mmol} \times 210,5 = 2,10 \text{ gram}$$

$$\text{Massa praktis} = 1,22 \text{ gram}$$

$$\% \text{ hasil} = \frac{1,22}{2,10} \times 100\% = 58,09 \% \approx 58 \%$$

## LAMPIRAN B

### UJI ANOVA 1

	Sum of Square s	Df	Mean Square	F	Sig.
Between Groups	82.667	2	41.333	74.400	.000
Within Groups	3.333	6	.556		
Total	86.000	8			

### Post Hoc Tests

#### Multiple Comparisons

Dependent Variable: Hasil  
Tukey HSD

(I) senyawa	(J) senyawa	Mean Difference (I-J)	Std. Error	Sig.	95% Confidence Interval	
		Lower Bound			Lower Bound	Upper Bound
1	2	-7.333(*)	.609	.000	-9.20	-5.47
	3	-4.667(*)	.609	.001	-6.53	-2.80
2	1	7.333(*)	.609	.000	5.47	9.20
	3	2.667(*)	.609	.011	.80	4.53
3	1	4.667(*)	.609	.001	2.80	6.53
	2	-2.667(*)	.609	.011	-4.53	-.80

\* The mean difference is significant at the .05 level.

## Homogeneous Subsets

### Hasil

Tukey HSD

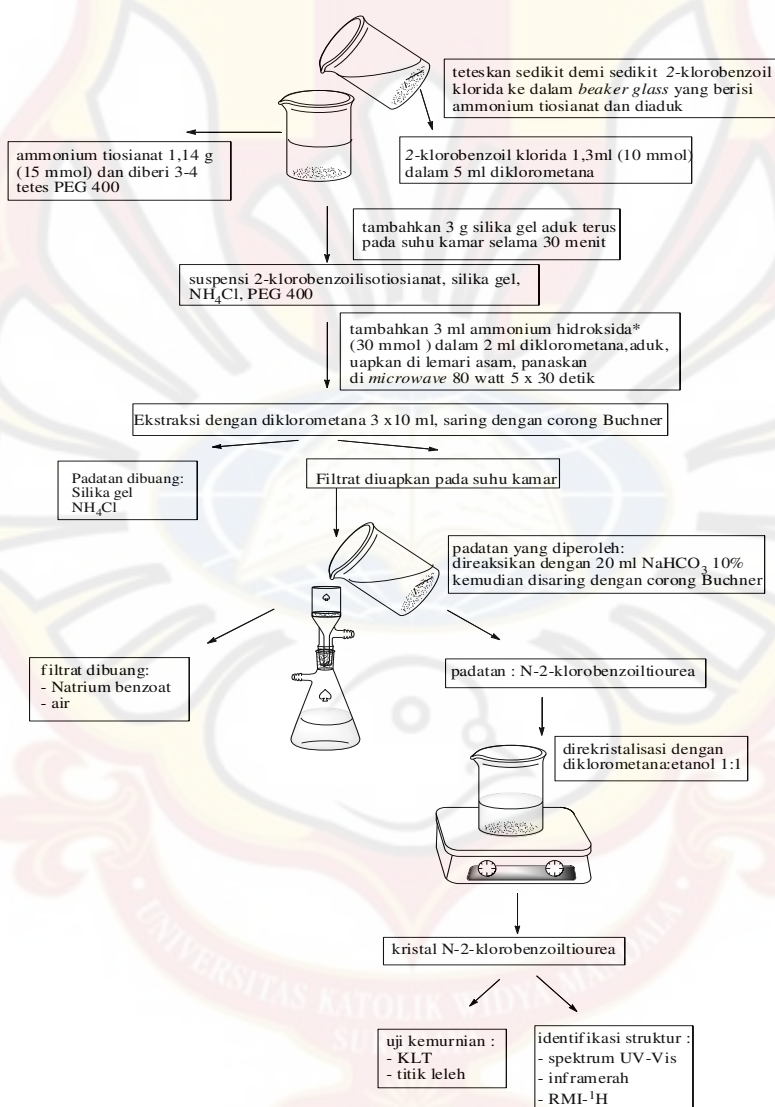
senyawa	N	Subset for alpha = .05		
		2	3	1
1	3	57.00		
3	3		61.67	
2	3			64.33
Sig.		1.000	1.000	1.000

Means for groups in homogeneous subsets are displayed.

a Uses Harmonic Mean Sample Size = 3.000.

## LAMPIRAN C

### SKEMA KERJA SINTESIS TURUNAN N-2-KLORO BENZOILTIOUREA



Keterangan \*:

Sintesis 2-kloro-N-(morpholin-4-karbonotiol)benzamida : 2.59 ml (30 mmol)

Morpholin

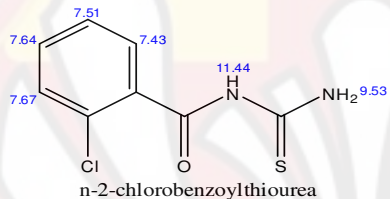
Sintesis 2-kloro-N-(piperazin-1-karbonotiol)benzamida : 2.58 g (30 mmol)

Piperazin

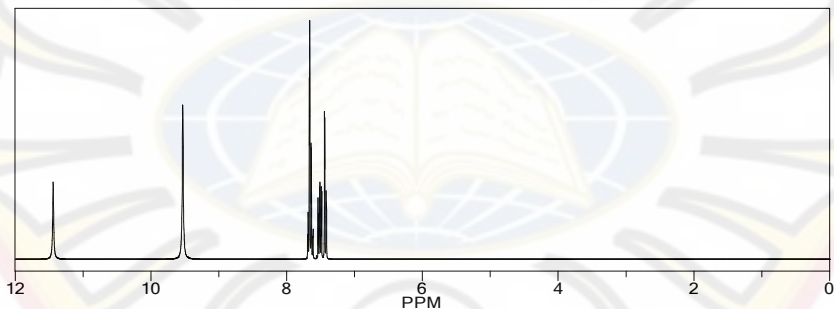
## LAMPIRAN D

### ESTIMASI RMI-<sup>1</sup>H SENYAWA N-2-KLOROBENZOILTIOUREA

#### ChemNMR <sup>1</sup>H Estimation



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



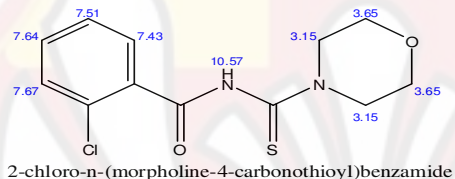
#### Protocol of the H-1 NMR Prediction:

Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
NH	11.44	8.00 3.44	sec. amide general corrections
NH2	9.53	2.00 7.53	amine general corrections
CH	7.67	7.26 0.01 0.18 0.22	1-benzene 1 -Cl 1 -C(=O) N general corrections
CH	7.43	7.26 -0.06 0.69 -0.46	1-benzene 1 -Cl 1 -C(=O) N general corrections
CH	7.64	7.26 -0.06 0.25 0.19	1-benzene 1 -Cl 1 -C(=O) N general corrections
CH	7.51	7.26 -0.12 0.18 0.19	1-benzene 1 -Cl 1 -C(=O) N general corrections

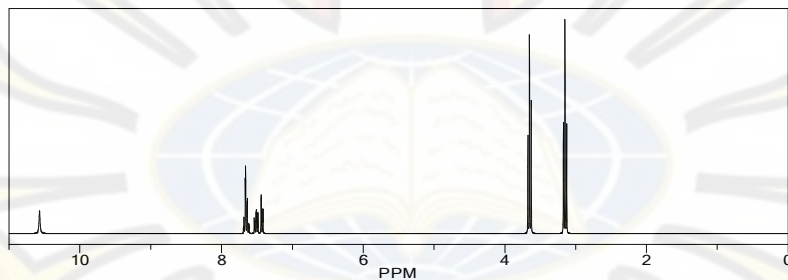
## LAMPIRAN E

### ESTIMASI RMI-<sup>1</sup>H SENYAWA 2-KLORO-N-(MORFOLIN-4-KARBONOTIOL)BENZAMIDA

#### ChemNMR <sup>1</sup>H Estimation



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



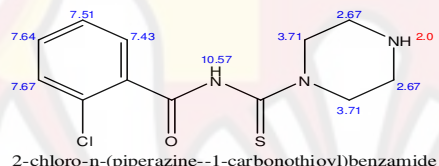
Protocol of the H-1 NMR Prediction:

Node	Shift	Base + Inc.	Comment (ppm rel. to TMS)
NH 10.57	8.00	sec. amide	
	2.57	general corrections	
CH2 3.65	3.67	tetrahydro-1,4-oxazine	
	-0.02	general corrections	
CH2 3.65	3.67	tetrahydro-1,4-oxazine	
	-0.02	general corrections	
CH2 3.15	2.87	tetrahydro-1,4-oxazine	
	?	1 -R from N-CHx	
	0.28	general corrections	
CH2 3.15	2.87	tetrahydro-1,4-oxazine	
	?	1 -R from N-CHx	
	0.28	general corrections	
CH 7.67	7.26	1-benzene	
	0.01	1 -Cl	
	0.18	1 -(C(=O)N	
	0.22	general corrections	
CH 7.43	7.26	1-benzene	
	-0.06	1 -Cl	
	0.69	1 -(C(=O)N	
	-0.46	general corrections	
CH 7.64	7.29	1-benzene	
	-0.06	1 -Cl	
	0.25	1 -(C(=O)N	
	0.19	general corrections	
CH 7.51	7.29	1-benzene	
	-0.12	1 -Cl	
	0.18	1 -(C(=O)N	
	0.19	general corrections	

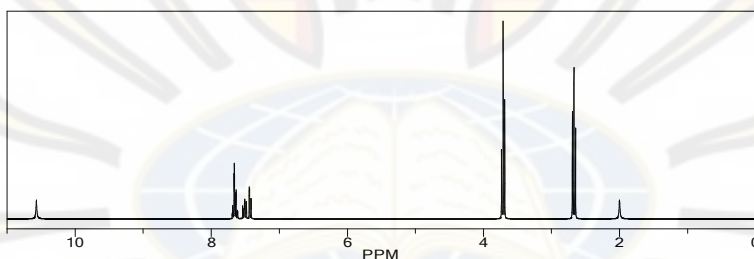
## LAMPIRAN F

### ESTIMASI RMI-<sup>1</sup>H SENYAWA 2-KLORO-N-(PIPERAZIN-1-KARBONOTIOL)BENZAMIDA

ChemNMR <sup>1</sup>H Estimation



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)
NH	2.0	2.00	amine
NH	10.57	8.00 2.57	sec. amide general corrections
CH2	3.71	1.37	methylene
		2.12 0.08 0.14	1' alpha -NC(=S)NR 1 beta -N-C general corrections
CH2	3.71	1.37	methylene
		2.12 0.08 0.14	1 alpha -NC(=S)NR 1 beta -N-C general corrections
CH2	2.67	1.37	methylene
		1.22 0.08	1 alpha -N-C 1 beta -N-C
CH2	2.67	1.37	methylene
		1.22 0.08	1 alpha -N-C 1 beta -N-C
CH	7.67	7.26	1-benzene
		0.01 0.18 0.22	1 -Cl 1 -(C=O)N general corrections
CH	7.43	7.26	1-benzene
		-0.06 0.69 -0.46	1 -Cl 1 -(C=O)N general corrections
CH	7.64	7.26	1-benzene
		-0.06 0.25 0.19	1 -Cl 1 -(C=O)N general corrections
CH	7.51	7.26	1-benzene
		-0.12 0.18 0.19	1 -Cl 1 -(C=O)N general corrections

