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INVESTIGATION OF REACTION OF VARIOUS THIOPHENE BASED KNOEVENAGEL ADDUCTS WITH ACETOACETANILIDE

F.N. Naghiyev, A.M. Maharramov, A.R. Asgarova, A.G. Rahimova,
M.A. Akhundova, I.G. Mamedov

Baku State University

23, Z.Khalilov str., AZ-1148 Baku, Azerbaijan; e-mail: farid.orgchemist@gmail.com

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The reaction of Michael addition interaction of 3-phenyl-2-(thiophene-2-carbonyl)acrylonitrile, 2-(thiophene-2-carbonyl)-3-(p-tolyl)acrylonitrile, 3-(4-methoxyphenyl)-2-(thiophene-2-carbonyl)acrylonitrile, as well as 3-pyridinyl-2-(thiophene-2-carbonyl)acrylonitrile with acetoacetanilide made it possible to produce substituted hexanones and 3,4-dihydro-2H-pyran derivatives. Structures of synthesized compounds were acknowledged by NMR and X-Ray structural analysis.

Keywords: thiophene, acrylonitrile, acetoacetanilide, pyran, NMR

INTRODUCTION

Pyranes and cyclic ketones are biologically active compounds to make up a structural part of natural compounds [1-2]. New [4+2] annulations were carried out in the presence of triphenylphosphine catalyst and high yield synthesized functionalized dihydropyranes [3]. A similar reaction of α -valin derived phosphine was used as catalyst in another work [4]. Also, new cyclation was carried out for α,β -unsaturated ketones through the use of DABCO-catalyst and synthesized

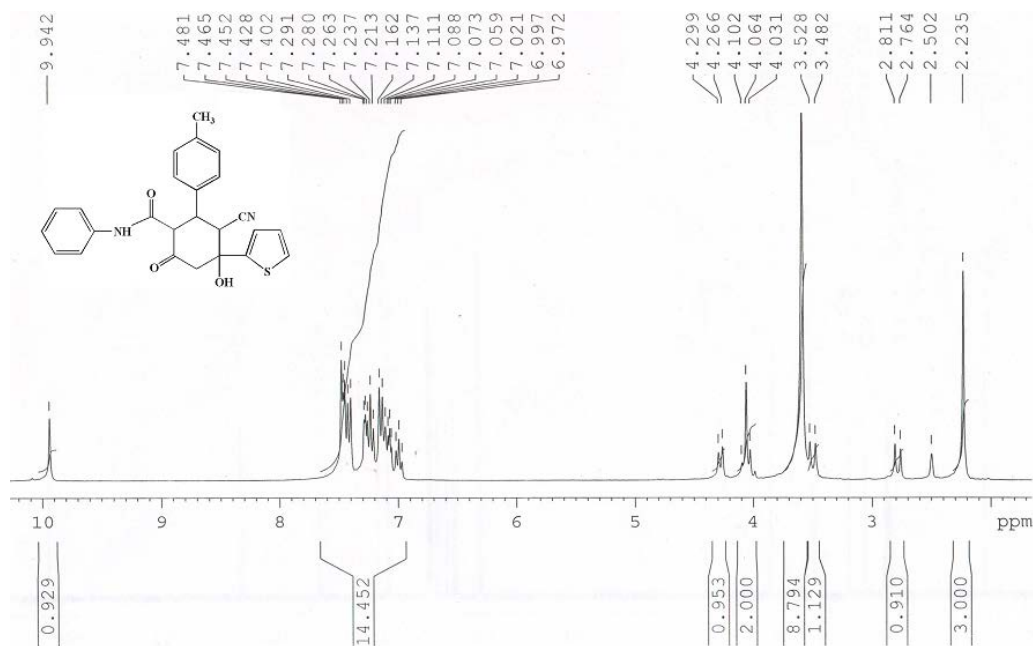
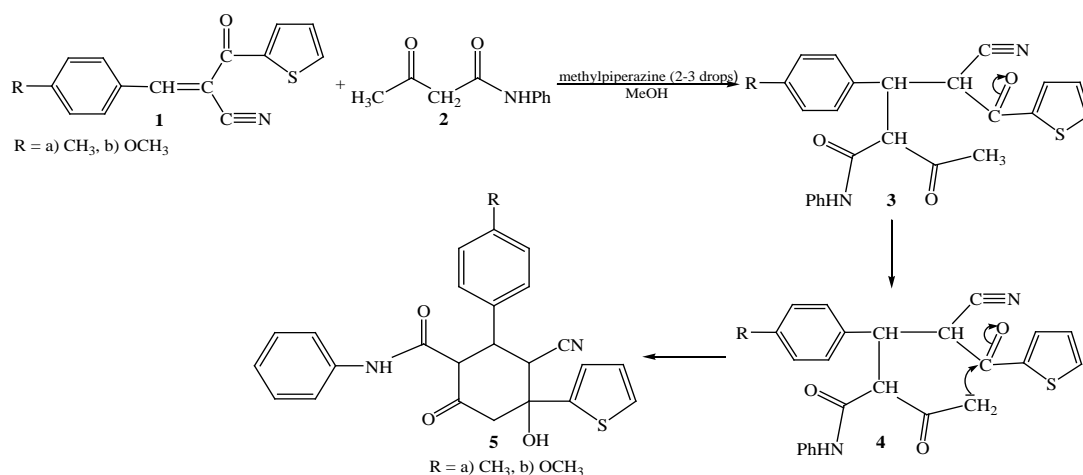
diastereomeric dihydropyran derivatives [5]. A cascade reaction of malononitrile and α -substituted calcone allowed to synthesize chiral multi-substituted amino-4H-pyran derivatives. Alkaloids were checked up as catalyst in pyran synthesis reactions [6].

The reaction of isatylidenmalononitriles with malononitriles and 2-thiophenemethylamine (or furfuryl amine) one-pot three component condensation was carried out for the first time to obtain appropriate spiroiridines [7].

RESULTS AND DISCUSSION

We carried out an experiment through the use of the Michael addition reaction for 2-(thiophene-2-carbonyl)-3-(p-tolyl)acrylonitrile and 3-(4-methoxyphenyl)-2-(thiophene-2-carbonyl)acrylonitrile with acetoacetanilide at room temperature in methanol media using of 2-3 drops of methylpiperazine to obtain corresponding hexanone-substituted derivatives. As

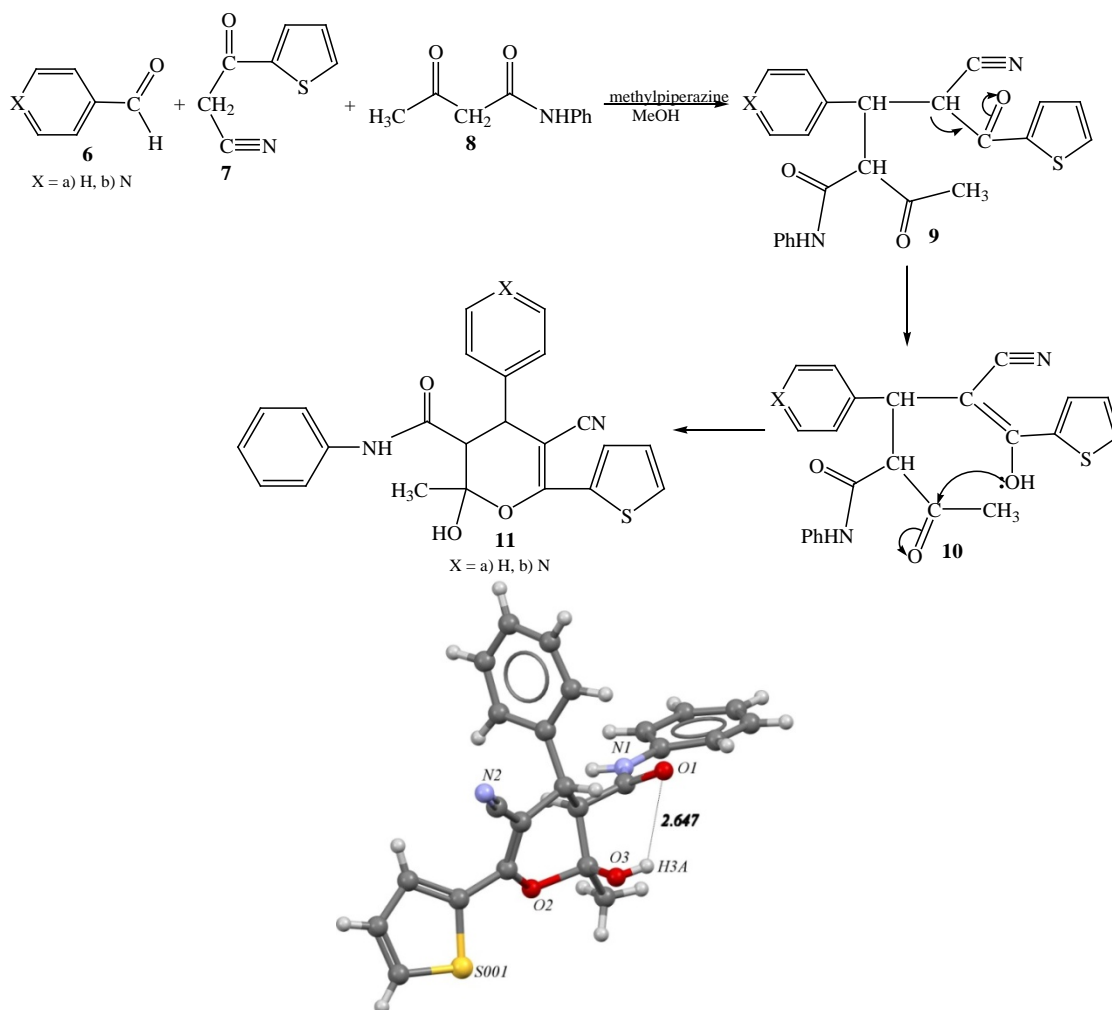
follows from the reaction mechanism, at the first stage the Michael addition of methylene active compound to double bond takes place and from the obtained Michael adduct (3) forms CH_2 -anion by action of base. At the last stage this anion attacks a carbonyl group of anion to synthesize an appropriate reaction products(5).



¹H NMR spectra of 3-cyano-4-hydroxy-6-oxo-N-phenyl-4-(thiophen-2-yl)-2-(p-tolyl)cyclohexane-1-carboxamide (**5a**)

Subsequently by means of the Michael addition reaction of Knoevenagel adduct obtained as a result of interaction between pyridine aldehyde and 2-thenoylacetonitrile,

with acetoacetanilide in ethanol-water media through the use of 2-3 drops of methylpiperazine to obtain appropriate 3,4-dihydro-2*H*-pyrane derivatives.



X-Ray structure of 5-cyano-2-hydroxy-2-methyl-N,4-diphenyl-6-(thiophen-2-yl)-3,4-dihydro-2H-pyran-3-carboxamide (**11a**)

EXPERIMENTAL PART

It should be noted that all used reagents were bought from Merc and Fluca and used without cleaning. The melting points of compounds were measured at STUART SPM30. Purity of synthesized compounds were checked by TLC, and structures acknowledged on "Bruker 300" NMR apparatus (300 and 75 MHz).

3-Cyano-4-hydroxy-6-oxo-N-phenyl-4-(thiophen-2-yl)-2-(p-tolyl)cyclohexane-1-carboxamide (5a): Mixture of 2-(thiophene-2-carbonyl)-3-(p-tolyl)acrylonitrile (2.9 mmol) and acetoacetanilide (3 mmol) dissolved in 35 ml of methanol was stirred for 5-7 minutes and a 2-3 drop methylpiperazine was added, so the stirring was continued. The reaction progress was tracked by TLC (EtOAc/n-hexane, 2:1) and the mixture kept quietly for

24-48 hours. Through evaporating of solvent it became possible to precipitate crystals. Note that crystals were separated by filter paper and recrystallized from ethanol (95%) - water mixture, yield 0.89 g, 71.60%. mp. 210°C.

¹H NMR (300 MHz, DMSO-*d*₆), δ, m.h.: 2.23 (s, 3H, CH₃); 2.79 (d, 1H, CH, ³J_{H-H} = 14.1); 3.50 (d, 1H, CH, ³J_{H-H} = 13.8); 4.06 (s, 2H, CH₂); 4.28 (d, 1H, CH, ³J_{H-H} = 9.9); 6.97-7.48 (m, 12H, 9Ar-H+3CH_{thiophenyl}); 9.94 (s, 1H, NH). ¹³C NMR spektr (DMSO-*d*₆), δ, m.h.: 21.14 (CH₃-Ar), 44.26 (CH), 47.40 (CH), 54.07 (CH₂), 62.64 (CH), 75.29 (C_{quat.}), 119.02 (CN), 119.49 (2CH_{arom}), 123.87 (CH_{thiophenyl}), 124.45 (CH_{arom}), 125.71 (CH_{thiophenyl}), 127.63 (CH_{thiophenyl}), 128.75 (2CH_{arom}), 129.14 (2CH_{arom}), 129.54 (2CH_{arom}), 137.06 (C_{ar}),

137.17 (C_{ar}), 139.14 (C_{ar}), 150.57 (C_{thioph}), 165.85 (CONH), 203.12 (C=O).

Found, %: 69.70 C; 5.07 H. C₂₅H₂₂N₂O₃S. Calculated, %: 69.77 C; 5.12 H. **3-Cyano-4-hydroxy-2-(4-methoxy-phenyl)-6-oxo-N-phenyl-4-(thiophen-2-yl)-cyclohexane-1-carboxamide (5b)**: it was synthesized in the same condition (yield 1.1 g, 85.61%). mp. 208°C.

¹H NMR (300 MHz, DMSO-*d*₆), δ, m.h.: 2.77 (d, 1H, CH, ³J_{H-H} = 14.1); 3.48 (d, 1H, CH, ³J_{H-H} = 15.3); 3.69 (s, 3H, CH₃O); 4.03 (s, 2H, CH₂); 4.27 (d, 1H, CH, ³J_{H-H} = 9.6); 6.89-7.47 (m, 12H, 9Ar-H+3CH_{thiophenyl}); 9.93 (s, 1H, NH). ¹³C NMR spektr (DMSO-*d*₆), δ, m.h.: 43.88 (CH), 47.51 (CH), 54.06 (CH₂), 55.37 (CH₃O), 62.79 (CH), 75.22 (C_{quat.}), 114.23 (2CH_{arom}), 119.07 (CN), 119.52 (2CH_{arom}), 123.87 (CH_{thiophenyl}), 124.42 (CH_{arom}), 125.70 (CH_{thiophenyl}), 127.62 (CH_{thiophenyl}), 129.14 (2CH_{arom}), 129.98 (2CH_{arom}), 131.96 (C_{ar}), 139.13 (C_{ar}), 150.59 (C_{thioph}), 158.90 (O-C_{ar}), 165.90 (CONH), 203.14 (C=O).

Found, %: 67.50 C; 4.45 H. C₂₅H₂₀N₂O₄S. Calculated, %: 67.57 C; 4.50 H. **5-Cyano-2-hydroxy-2-methyl-N,4-diphenyl-6-(thiophen-2-yl)-3,4-dihydro-2H-pyran-3-carboxamide (11a)**:

Mixture of 3-phenyl-2-(thiophene-2-carbonyl)acrylonitrile (2.9 mmol) and acetoacetanilide (3 mmol) dissolved in 35 ml of methanol was stirred for 5-7 minutes and a 2-3 drop methylpiperazine was added, so the stirring was continued. Reaction progress was tracked by TLC (EtOAc/n-hexane, 2:1), and the mixture was kept quietly for 24-48 hours. By evaporating of solvent crystals were precipitated. Crystals were separated by filter paper and recrystallized from ethanol (95%) - water mixture. yield 0.9 g, 75%. mp. 174°C.

¹H NMR (300 MHz, DMSO-*d*₆), δ, m.h.: 1.72 (s, 3H, CH₃); 3.07 (d, 1H, CH, ³J_{H-H} = 11.7); 4.38 (d, 1H, CH, ³J_{H-H} = 11.7); 7.00-7.89 (m, 13H, 10Ar-H+3CH_{thiophenyl}); 9.86 (s, 1H, NH). ¹³C NMR spektr (DMSO-*d*₆), δ,

m.h.: 26.38 (CH₃), 40.62 (CH), 55.87 (CH), 86.25 (=C_{quat.}), 99.73 (O-C_{quat.}), 119.58 (CN), 119.66 (2CH_{arom}), 124.15 (CH_{arom}), 128.03 (CH_{arom}), 128.47 (CH_{thiophenyl}), 128.60 (CH_{arom}), 128.87 (2CH_{arom}), 129.05 (3CH_{arom}), 129.78 (CH_{thiophenyl}), 130.63 (CH_{thiophenyl}), 135.99 (C_{ar}), 138.87 (C_{thioph}), 140.12 (C_{ar}), 166.77 (CONH), 167.63 (O-C_{quat.}=).

Found, %: 69.17 C; 4.76 H. C₂₄H₂₀N₂O₃S. Calculated, %: 69.23 C; 4.81 H.

5-Cyano-2-hydroxy-2-methyl-N-phenyl-4-(pyridin-4-yl)-6-(thiophen-2-yl)-3,4-dihydro-2H-pyran-3-carboxamide (11b):

Mixture 4-pyridinecarboxaldehyde or (2.9 mmol) and 2-thenoylacetonitrile (3 mmol) dissolved in 50 ml of ethanol-water (4:1) was stirred for 5 minutes, reaction mixture was kept quietly for 36 hours. Then acetoacetanilide (3 mmol) was added to reaction mixture, stirred for 5 minutes and after 2-3 drop methylpiperazine was added and stirring was continued. Reaction progress was tracked by TLC (EtOAc/n-hexane, 2:1). Reaction mixture was kept quietly for 36 hours. Through evaporating of solvent it became possible to precipitate crystals. Crystals were separated by filter paper and recrystallized from ethanol (95%) - water mixture. yield 0.97 g, 79.51%. mp. 165°C.

¹H NMR (300 MHz, DMSO-*d*₆), δ, m.h.: ¹³C NMR spektr (DMSO-*d*₆), δ, m.h.: 1.70 (s, 3H, CH₃); 3.05 (d, 1H, CH, ³J_{H-H} = 11.4); 4.39 (d, 1H, CH, ³J_{H-H} = 11.4); 7.03-8.56 (m, 12H, 9Ar-H+3CH_{thiophenyl}); 9.88 (s, 1H, NH). ¹³C NMR spektr (DMSO-*d*₆), δ, m.h.: 26.31 (CH₃), 40.21 (CH), 55.04 (CH), 84.45 (=C_{quat.}), 99.72 (O-C_{quat.}), 119.71 (CN), 120.02 (2CH_{arom}), 124.29 (CH_{arom}), 124.44 (CH_{arom}), 128.52 (CH_{thiophenyl}), 129.21 (3CH_{arom}), 130.11 (CH_{thiophenyl}), 130.97 (CH_{thiophenyl}), 135.67 (C_{ar}), 138.68 (C_{thioph}), 149.17 (C_{ar}), 150.38 (2CH_{arom}), 166.68 (CONH), 167.23 (O-C_{quat.}=).

Found, %: 66.12 C; 4.51 H. C₂₃H₁₉N₃O₃S. Calculated, %: 66.19 C; 4.56 H.

REFERENCES

1. Aleksandra Pałasz. Synthesis of 3,4-dihydro-2H-pyrans by hetero-Diels–Alder reactions of functionalized α,β -unsaturated carbonyl compounds with N-vinyl-2-oxazolidinone. *Org. Biomol. Chem.*, 2005, vol.3, iss.17, pp. 3207-3212. doi: 10.1039/B504210K
2. Dhananjay B. Kendre, Raghunath B. Toche, Madhukar N. Jachak. Michael addition of dimedone with α,β -unsaturated ketones: Synthesis of quinoline and chromene derivatives. *Journal of Heterocyclic Compounds*. 2008, vol. 45, iss. 3, pp. 667-671. doi: org/10.1002/jhet.5570450305
3. Qiongmei Zhang, Tong Fang, Xiaofeng Tong. Facile synthesis of highly functionalized six-membered heterocycles via PPh_3 -catalyzed [4+2] annulations of activated terminal alkynes and heterodienes: scope, mechanism, and application. *Tetrahedron*. 2010, vol. 66, iss. 40, pp. 8095-8100. doi: org/10.1016/j.tet.2010.07.043
4. Huanzhen Ni, Weijun Yao, Abdul Waheed, Nisar Ullah, and Yixin Lu. Enantioselective [4+2]-Annulation of Oxadienes and Allenones Catalyzed by an Amino Acid Derived Phosphine: Synthesis of Functionalized Dihydropyrans. *Org. Lett.*, 2016, vol.18, iss. 9, pp.2138–2141. doi: 10.1021/acs.orglett.6b00760
5. Wen Liu, Jing Zhou, Changwu Zheng, Xingkuan Chen, Hua Xiao, Yingquan Yang, Yinlong Guo, Gang Zhao. Tandem cross-RauhuteCurrier/cyclization reactions of activated alkenes to give densely functionalized 3,4-dihydropyrans. *Tetrahedron*. 2011, vol. 67, iss.10, pp.1768-1773. doi: org/10.1016/j.tet.2011.01.036
6. Zhi-Peng Hu, Wei-Juan Wang, Xiao-Gang Yin, Xue-Jing Zhang, Ming Yan. Enantioselective synthesis of 2-amino-4H-pyrans via the organocatalytic cascade reaction of malononitrile and α -substituted chalcones. *Tetrahedron: Asymmetry*. 2012, vol. 23, iss. 6–7, pp. 461-467. doi.org/10.1016/j.tetasy.2012.03.018
7. Maharramov A.M., Naghiyev F.N., Asgarova A.R., Rahimova A.G., Akhundova M.A., Mamedov I.G. Investigation of conversion various ilidenmalononitriles. *Kimya Problemleri – Chemical Problems*. 2018, no.1, pp. 69-72. (In Azerbaijan).

**TİOFEN ƏSASLI BƏZİ KNOEVENAGEL ADDUKTLARININ ASETOASETANİLİDLƏ
REAKSİYASININ TƏDQIQI**

**F.N. Nağıyev, A.M.Məhərramov, A.R. Əsgərova, A.G. Rəhimova,
M.A. Axundova, I.G. Mamedov**

Bakı Dövlət Universiteti

AZ 1148 Bakı, Z.Xəlilov küç., 23; e-mail: farid.orgchemist@gmail.com

3-Fenil-2-(tiofen-2-karbonil)akrilonitril, 2-(tiofen-2-karbonil)-3-(p-tolil)akrilonitril və 3-(4-metoksifenil)-2-(tiofen-2-karbonil)akrilonitril, eləcə də 3-piridinil-2-(tiofen-2-karbonil)akrilonitrilin asetoasetanilid ilə Mixael birləşmə reaksiyasıyından əvəzlənmiş heksanon və 3,4-dihidro-2H-piran törəmələrinin alınması müəyyən edilmişdir. Sintez edilən birləşmələrin quruluşu NMR və RQA analiz metodlarının köməyiylə təsdiq olunmuşdur.

Açar sözlər: tiofen, akrilonitril, asetoasetanilid, piran, NMR

**ИССЛЕДОВАНИЕ РЕАКЦИИ ВЗАИМОДЕЙСТВИЯ НЕКОТОРЫХ АДДУКТОВ
КНЕВЕНАГЕЛЯ НА ОСНОВЕ ТИОФЕНА С АЦЕТОАЦЕТАНИЛИДОМ**

**Ф.Н. Нагиев, А.М. Магеррамов, А.Р. Аскерова, А.Г. Рагимова,
М.А. Ахундова, И.Г. Мамедов**

*Бакинский государственный университет
AZ 1148 Баку, ул. З.Халилова, 23; e-mail: farid.orgchemist@gmail.com*

Путем реакции присоединения по Михаэлю взаимодействием 3-фенил-2-(тиофен-2-карбонил) акрилонитрила, 2-(тиофен-2-карбонил)-3-(n-толил)акрилонитрила и 3-(4-метоксифенил)-2-(тиофен-2-карбонил)акрилонитрила, а также 3-пиридинил-2-(тиофен-2-карбонил)акрилонитрила с ацетоацетанилидом были получены соответствующие замещенные гексаноны и 3,4-дигидро-2H-пиран-производные. Структуры синтезированных соединений были подтверждены методами ЯМР и рентгеноструктурного анализа.

Ключевые слова: *тиофен, акрилонитрил, ацетоацетанилид, пиран, ЯМР*