**Full Paper** 

Reference Object Identifier - ROI: jbc-02/18-54-4-88 Publication is available for discussion in the framework of the on-line Internet conference "Butlerov readings". http://butlerov.com/readings/ Submitted on March 15, 2018.

## Synthesis of di- and trialkenylsubstituted aromatic amines

© Akhat G. Mustafin,\*<sup>1,2</sup> Ildus B. Abdrakhmanov,<sup>2</sup> Vakil M. Sharafutdinov,<sup>3</sup> and Yury S. Zimin<sup>1+</sup>

<sup>1</sup>Department of Physical Chemistry and Chemical Ecology. Bashkir State University. Zaki Validi St., 32. Ufa, 450074. Republic of Bashkortostan. Russia. Phone: +7 (347) 229-96-94. E-mail: ZiminYuS@mail.ru <sup>2</sup> Laboratory of Pharmacophore Cyclic Systems. Ufa Institute of Chemistry – Subdivision of the Ufa Federal Research Centre of the Russian Academy of Sciences. Oktyabrya Ave., 71. Ufa, 450054. Republic of Bashkortostan. Russia. <sup>3</sup> Department of General Chemical Technology. Branch of Ufa State Petroleum Technological University

in Sterlitamak. Oktyabrya Ave., 2. Sterlitamak, 453118. Republic of Bashkortostan. Russia.

\*Supervising author; <sup>+</sup>Corresponding author

Keywords: aromatic amines, allyl chloride, 4-chloro-2-pentene, alkenylation, rearrangement of Claisen.

## Abstract

Inventing the Claisen rearrangement in aromatic amines series has led to bio-active 2-alkenylsubstituted arylamines availability and their wide usage in the synthesis of heterocyclic compounds. The presence of several reaction centers in the allyl aniline molecule makes it possible to carry out their further transformations in the nitrogen atom, the alkenyl unit, or two centers simultaneously. The preparative value of these compounds can be further strengthened by adding active functional groups or a double bond to the molecule. In this paper, the possibility of introducing an alkenyl group into the composition of aromatic amines with an allyl-type substituent in the nucleus or at the nitrogen atom has been considered. Continuous boiling (20-30 h) of 2-(1-methyl-2-butenyl)arylamines with 4-chloro-2-pentene in a triethylamine solution leads to their N-substituted derivatives. The same reaction with 3-chloropropene and 3-chloro-2methylpropene is completed in 8 hours. The introduction of the second alkenyl group along the nitrogen atom of N-(1-methyl-2-butenyl)aniline is under harder conditions (150 °C, autoclave), with 4-chloro-2-pentene forming N,2-di-(1-methyl-2-butenyl)aniline (I). By heating I under ZnCl<sub>2</sub> in nitrobenzene, 2,6-di-(1-methyl-2butenyl)aniline (II) with 78% yield and a small amount of 2,4,6-tri-(1-methyl-2-butenyl)aniline (III) were obtained. The same products were obtained by direct 4-chloro-2-pentene and a twofold excess of 2-(1-methyl-2-butenyl)aniline reaction with 10% III yield. Both schemes have been implemented to get 2,6-di-(1-methyl-2-butenyl)-4-methylaniline (IV). N,2-Di-(1-methyl-2-butenyl)-4-methylaniline prepared by the method described, is rearranged under ZnCl<sub>2</sub> in IV with 82% yield. The same result was obtained by heating 4-chloro-2-pentene at 150 °C with a twofold excess of 2-(1-methyl-2-butenyl)-4-methylaniline. To synthesize 2,4-di-(1-methyl-2-butenyl)-6-methylaniline (V) according to the scheme developed, N.2-di-(1-methyl-2-butenyl)-6methylaniline (VI) under equimolar amount of ZnCl<sub>2</sub> in nitrobenzene leads to the amine V desired. The catalysis of the Claisen VI rearrangement with an equimolar amount of ortho-toluidine hydrochloride at 140-150 °C also leads to V, but the yield is low, since the splitting process predominates under these conditions. Direct reaction II with 4-chloro-2-pentene is not feasible due to the steric inhibition of the N-alkenylation step by volume substituents in the *ortho*-positions. The structure of poly-substituted arylamines is established by spectral methods and elemental analysis. The compounds obtained are of interest for the nitrogen-containing heterocycle synthesis.

## References

- [1] I.B. Abdrakhmanov, A.G. Mustafin, V.M. Sharafutdinov. Rearrangement of Claisen in a row of aromatic amines. Ufa: Gilem, Bashkir encvclopedia. 2014. 168p. (russian)
- C.D. Hurd, W.W. Jenkins. Decomposition of allylanilines with the presence of zinc chloride. Journal of [2] Organic Chemistry. 1957. Vol.22. P.1418-1423.
- [3] R.R. Gataullin. Preparation a Heterocyclic Compounds from the N- and C-alkenylanilines. Vestnik of St. Petersburg State University. 2014. Series 4. Iss.1. P.51-70. (russian)

## SYNTHESIS OF DI- AND TRIALKENYLSUBSTITUTED AROMATIC AMINES

- [4] I.B. Abdrakhmanov, V.M. Sharafutdinov, N.G. Nigmatullin, I.A. Sagitdinov, G.A. Tolstikov. Amino-Claisen Rearrangement as a Method of Synthesis of C-substituted Amines. Russian Journal of Organic Chemistry. 1982. Vol.18. No.7. C.1466-1471. (russian)
- [5] R.R. Gataullin, T.V. Kazhanova, I.A. Sagitdinov, A.A. Galyautdinov, A.A. Fatykhov, L.V. Spirikhin, I.B. Abdrakhmanov. Alkenylation of anilines with dicyclopentadiene, cyclopentadiene and piperylen. Russian Journal of Applied Chemistry. 2001. Vol.74. Iss.2. P.274-280. (russian)
- [6] R.N. Khusnitdinov, A.N. Khazipova, K.R. Khusnitdinov, B.I. Kutepov, I.B. Abdrakhmanov, A.G. Mustafin. Amino-Claisen Rearrangement of N-(1'-methyl-2'-buthenyl)aniline at the Presence of Different Catalysts. Bashkir Chemical Journal. 2014. Vol.21. No.1. P.87-91. (russian)
- [7] I.B. Abdrakhmanov, V.M. Sharafutdinov, U.M. Dzhemilev, E.V. Talvinskiy, I.A. Sagitdinov, G.A. Tolstikov. Synthesis of N-substituted Unsaturated Amines. Russian Journal of Applied Chemistry. 1982. No.9. P.2121-2123. (russian)