Multicomponent Synthesis of 2, 4, 6-Triarylpyridine Analogues As Antibacterial And Antiurease Agents

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Abstract

Five new 2,4,6-triarylpyridine analogues with potential biological importance have been synthesized using the multicomponent reaction (MCR) solid support catalyst synthetic approach. This green synthesis furnished method simplicity, reduce reaction time, and excellent yield, along with an additional feature of catalyst reusability. A number of spectroscopic techniques such as $^1$H-NMR, IR, and mass spectrometry, confirmed structures of the newly synthesized compounds. Antibacterial and anti-urease activities of these compounds were evaluated. Results revealed that compounds 4a, 4c, 4d, and 4e exhibit significant inhibition against gram-positive bacterial pathogens. Furthermore, all synthesized compounds showed potent urease inhibitory activity with IC$_{50}$ values ranging from 12.8 ± 1.04 to 23.7 ± 0.23 µM when compared with the standard inhibitor thiourea (IC$_{50}$ 21.0 ± 0.23 µM).

In addition, the synthesized derivatives were subjected to density functional theory (DFT) calculations (DFT) to determine the energy of the molecules having biological potential.

Introduction

Microbial and viral infections continue to be a major health problem and a great challenge worldwide due to their potential resistance. Medicinal chemists have been persistently struggling to develop novel and effective antibiotics to overcome this problem (Mouradzadegun, Elahi, Ghanbarzadeh, Silicon, & Elements, 2015). In this context, pyridine analogues have received considerable attention due to their unique structural features and diverse biological properties, which range from antibacterial, to antitumor (Ren & Cai, 2009). Examples of such compounds that are marketed pharmaceuticals include esomeprazole, used to treat symptoms of gastroesophageal reflux disease (GERD), and can lower the risk of bleeding after endoscopy in patients with ulcers. Other examples include rimiterol and etoricoxb (Cotton et al., 2000). Rimiterol is a direct-acting sympathomimetic with predominantly β-adrenergic activity and a selective action on β2-receptors; it is a third-generation short-acting β$\text{\textregistered}$ agonist. Etoricoxb, on the other hand, is a selective inhibitor of cyclooxygenase-2 (COX-2), an enzyme involved in pain and inflammation, and is a nonsteroidal anti-inflammatory drugs (NSAIDs). Among other classes, 2,4,6-triarylpyridines are of great interest due to their wide range of biological and therapeutic potential.

Published research indicated that a large variety of catalysts such as heteropolyacid H$_{14}$[NaP$_5$W$_{30}$O$_{110}$], Fe$_3$O$_4$ supported silica sulfuric acid nanoparticles, BaCl$_2$-nano SiO$_2$, nano-ZnO powder, magnetic MIL-101-SO$_3$H, heterogeneous silica vanadic acid [SiO$_2$-VO(OH)$_2$], montmorillonite K10 clay, nanocrystals MgAl$_2$O$_4$, Brønsted-acidic ionic liquid [HO$_3$S(CH$_2$)$_4$MIM][HSO$_4$], PEG$_{1000}$ DAIL, acidified titanium silicates have been used in one-pot synthesis of 2,4,6-triarylpyridine analogues (Alinezhad, Tajbakhsh, & Ghibadi, 2015; Boroujeni, Hashemzadeh, Faroughi, Shaabani, & Amini, 2016; Davoodnia, Bakavoli, Moloudi, Tavakoli-Hoseini, & Khashi, 2010; Gadekar & Lande, 2018; Heravi, Bakhtiar, Daroogheha, & Bamoharram, 2007; Kannan & Sreekumar, 2013; Ren, Zhang, & Jin, 2016; Reza, Shaee, Moloudi, & Ghashang, 2012; Safari, Gandomi-Ravandi, & Boroujeni, 2013; Shafiee & Moloudi, 2011; Zolfigol et al., 2015). However, these
syntheses suffer from a number of shortcomings including the use of expensive catalysts, and the removal of catalyst and additional purification steps, thus compromising green chemistry principles. Another challenge is to construct multifunctional structures with linear approach that suffers from overall low yield. To overcome these problems, the one-pot MCR approach has been adopted to afford structural complexity and diversity in a single step which is cost effective (Sunderhaus & Martin, 2009). Accordingly, we have used the MCR method under solvent-free conditions, which is ecofriendly, and using a solid support silica impregnated with acid catalyst in the synthesis of new triarylpyridine analogues.

On the other hand, quantum chemical calculations are fascinating tools in chemistry as they rationalize the properties of newly synthesized compounds (A. J. C. m. s. Irfan, 2014). In this respect, the density functional theory (DFT) was comprehensively used to engross the geometric and electronic properties, which successfully reproduce experimental evidence (A. Irfan, Al-Sehemi, Chaudhry, & Muhammad, 2018). In addition, it is a reliable tool for the ground state (S₀) geometries and electronic properties (A. J. C. Irfan & Chemistry, 2019; Lee, Yang, & Parr, 1988). Similarly, B3LYP is rational for the S₀ geometries of various organic compounds (Preat, Jacquemin, Perpète, & technology, 2010; Preat, Michaux, Jacquemin, & Perpete, 2009). In the present study, optimizations were performed by B3LYP functional and triple zeta with 2 polarization function (TZ2P) basis sets. All calculations in this work were conducted using the Amsterdam Density Functional (ADF) modeling suite. Accordingly, we describe herein the synthesis and characterization of these new compounds (Schemes 1), along with the evaluation of their antibacterial and anti-urease activities. To the best of our knowledge, no previous study of this kind has appeared in the literature.

**Experimental**

**Materials and Methods**

All chemicals used throughout this investigation were purchased from Sigma Aldrich, Fluka and Merck, and were used as received. Progress of reaction and purity of products were monitored by thin layer chromatography (TLC) carried out on packed on silica gel Al (aluminum) sheets (60 F254). Plates were visualized under UV light, where appropriate. Catalyst was characterized by means of a scanning electron microscope (SEM), with accelerating voltage 0.2–30 kV of 50/60 Hz power by (Carl Zeiss Company). Infrared (FTIR) spectra were recorded as KBr pellets in the range of 650-4000 cm⁻¹ with a Shimadzu Corporation. ¹H NMR spectra of prepared compounds were recorded with the aid of a 500 MHz spectrometer (AVANCE AV). Chemical shifts are reported in ppm using tetramethylsilane (TMS) as an internal standard and CDCl₃ as solvent. (¹H NMR spectra of prepared compounds are displayed in Fig. 5S-Supplimentary).

¹H NMR spectra of prepared compounds are displayed in the supplementary material.

**Catalyst preparation**
Catalyst was prepared according to a published procedure which involved stirring a mixture of conc. sulfuric acid and silica gel in dichloromethane for 4 h; this has afforded a white amorphous powder, which was used as a solid support medium without further purification (Riego, Sedin, Zaldívar, Marziano, & Tortato, 1996).

**General procedure for preparation of 4a–e**

Compounds 4a–e were prepared by the following procedure: Equimolar amounts of arylaldehydes \( 1a–e \), 2-acetylfuran (2), 2-acetylthiophene (3), \( \text{NH}_4\text{OAc} \) (1.5 mole %), and an appropriate amount of silica catalyst were mixed in a round bottom flask and refluxed at 80 °C for 1 h (Table 1). Progress of reaction was monitored by TLC using (10 %) a mixture of \( n \)-hexane and ethyl acetate as a solvent system. Upon completion of reaction, the mixture was allowed to cool to room temperature, dissolved in methanol, and filtered. The filtrate was poured onto crushed ice to afford the final products (4a–e). Using the same general procedure, the following compounds were prepared:

**2-(Furan-2-yl)-4-phenyl-6-(thiophen-2-yl) pyridine (4a)**

Prepared as light brown sticky solid, yield 60 %; mp 113 °C; \( R_f = 0.72 \text{ cm}^{-1} \); IR (KBr) cm\(^{-1}\): 1559 (C=C), 1493 (C=N-C), 1235 (C-O-C), 1161 (C-S-C); \( ^1\text{H-NMR} \) (500 MHz, CDCl\(_3\)) \( \delta \) (ppm): 7.85 (1H, d, \( J = 2.1 \text{ Hz}, \text{H-3 pyridyl} \)), 7.82(1H, d, \( J = 2.1 \text{ Hz}, \text{H-5 pyridyl} \)), 7.62–7.78 (3H, m, Ar-H furfuryl), 7.40–7.32 (3H, m, Ar-H thienyl), 6.55–6.52 (5H, m, Ph).; MS \( m/z \) (ESI-MS) 304 (M\(^+\)) (90 %).

**2-(2-(Furan-2-yl)-6-(thiophen-2-yl) pyridin-4-yl) phenol (4b)**

Dark brown, gummy solid; yield 65 %; mp 112° C; \( R_f = 0.846 \text{ cm}^{-1} \); IR (KBr) cm\(^{-1}\): 1559 (C=C), 1510 (C=N-C), 1243 (C-O-C), 1157 (C-S-C); \( ^1\text{H-NMR} \) (500 MHz, CDCl\(_3\)) \( \delta \) (ppm):

7.74 (1H, d, \( J = 2.2 \text{ Hz}, \text{H-3 pyridyl} \)) 7.63 (1H, d, \( J = 2.3 \text{ Hz}, \text{H-5 pyridyl} \)), 7.61–7.52 (3H, m, Ar-H furfuryl), 7.35–7.31 (3H, m, Ar-H thienyl) 6.99 (1H, dd, \( J = 8.2, 2.1 \text{ Hz}, \text{Ph H-3} \)), 6.57 (1H, ddd, \( J = 8.3, 8.3, 2.1 \text{Hz}, \text{Ph H-5} \)), 6.55 (1H, dd, \( J = 8.2, 8.1 \text{ Hz}, \text{Ph H-4} \)), 6.53 (1H, dd, \( J = 8.2, 2.1 \text{ Hz}, \text{Ph H-6} \)).; MS \( m/z \) (ESI-MS) 320 (M\(^+\)) (90 %).

**2-(Furan-2-yl)-4-(4-methoxyphenyl)-6-(thiophen-2-yl) pyridine (4c)**

Light brown, sticky solid; yield 63 %; M.P. 110° C; \( R_f = 0.57 \text{ cm}^{-1} \); IR (KBr) cm\(^{-1}\): 1559 (C=C), 1510 (C=N-C), 1243 (C-O-C), 1172 (C-S-C); \( ^1\text{H-NMR} \) (500 MHz, CDCl\(_3\)) \( \delta \) (ppm):

7.85 (1H, d, \( J = 2.2 \text{ Hz}, \text{H-3 pyridyl} \)), 7.83 (1H, d, \( J = 2.2 \text{ Hz}, \text{H-5 pyridyl} \)), 7.62–7.59 (3H, m, Ar-H furfuryl), 7.32–7.28 (3H, m, Ar-H thienyl), 6.92 (2H, d, \( J = 8.2 \text{ Hz}, \text{H-3 and H-5} \)), 6.86 (2H, d, \( J = 8.2 \text{ Hz}, \text{H-2 and H-6} \)), 3.86 (3H, s, OCH\(_3\)) ; MS \( m/z \) (ESI-MS) 334 (M\(^+\)) (100 %).

**2,4-Di(furan-2-yl)-6-(thiophen-2-yl) pyridine (4d)**
Dark brown, sticky solid; yield 60 %; M.P. 145° C; R\textsubscript{f} = 0.57 cm\textsuperscript{-1}; IR (KBr) cm\textsuperscript{-1}: 1559 (C=C), 1507 (C=N-C), 1237 (C-O-C), 1156 (C-S-C); \textsuperscript{1}H-NMR (500 MHz, CDCl\textsubscript{3}) \delta (ppm): 7.62 (1H, d, J = 2.2 Hz, H-3 pyridyl), 7.60 (1H, d, J = 2.2 Hz, H-5 pyridyl), 7.79–7.73 (3H, m, Ar-H furfuryl), 7.32–7.29 (3H, m, Ar-H thiienyl), 6.96–6.70 (3H, m, Ar-H furfuryl); MS m/z (ESI-MS) 294 (M\textsuperscript{+}) (99 %).

\textbf{4-(4-Chlorophenyl)-2-(furan-2-yl)-6-(thiophen-2-yl) pyridine (4e)}

Light brown, granular solid; yield 55 %; M.P. 120° C; R\textsubscript{f} = 0.65 cm\textsuperscript{-1}; IR (KBr) cm\textsuperscript{-1}: 1559 (C=C), 1490 (C=N-C), 1247 (C-O-C), 1163 (C-S-C); \textsuperscript{1}H-NMR (500 MHz, CDCl\textsubscript{3}) \delta (ppm):

7.81 (1H, d, J = 2.2 Hz, H-3 pyridyl), 7.78 (1H, d, J = 2.2 Hz, H-5 pyridyl), 7.64–7.57 (3H, m, Ar-H furfuryl), 7.42–7.35 (3H, m, Ar-H thiienyl), 6.59 (2H, d, J = 8.2 Hz, H-3 and H-5), 6.38(2H, d, J = 8.2 Hz, H-2 and H-6); MS m/z (ESI-MS) 338 (M\textsuperscript{+}) (25 %).

\textbf{Biological Activities}

\textbf{In Vitro Cytotoxicity Assay}

Cytotoxicity of the newly synthesized 2,4,6-triarylpyridine derivatives has been performed in 96-well microtiter plates using the 3-[4, 5-dimethylthiazole-2-yl]-2,5-diphenyl-tetrazolium bromide (MTT) colorimetric assay. In this assay, mouse fibroblast (3T3) cells were cultured in Dulbecco’s Modified Eagle Medium, and mixed with 10 % of fetal bovine serum (FBS), 100 IU/mL of penicillin, and 100 1g/mL of streptomycin in 75 cm\textsuperscript{2} flask. Flasks were then kept in an incubator with 5% CO\textsubscript{2} at 37 °C. Cultured cells with a concentration of 5, 9, and 104 cells/mL were introduced (100 µL/well) into 96 well plates. After an incubation period of 12 h, 200 µL of fresh medium was added with different concentrations of compounds 4a–e (1 M). After 2 days, 200 µL of MTT (0.5 mg/mL) was added to each well, and kept incubated for 4 h. This was followed by addition of 100 µL of DMSO to each well. The degree of reduction of MTT to form a zone within cells was measured at 540 nm using a microplate reader. Cytotoxicity profile was calculated as the concentration causing 50% growth inhibition (IC\textsubscript{50}) of 3T3 cells (Saleem et al., 2016). The percent inhibition was calculated by using the following formula:

\[
\% \text{ inhibition} = 100 - \frac{\text{mean of OD test} - \text{mean of OD of negative control}}{\text{mean of OD positive control} - \text{mean of negative control}} \times 100
\]

\textbf{Antibacterial Screening}

We evaluated the antibacterial activity of synthesized compounds 4a–e using the disc diffusion method. Briefly, test compounds (10 mg/mL) were applied on the pasteurized filter paper discs, dried overnight at room temperature, and stored; the negative control discs were prepared using the same procedure. In a standard microbiological working environment, bacterial cultures were grown overnight at 37 °C in
nutrient broth medium and spread onto solidified nutrient agar medium in petri dishes. Then control and test discs were applied onto the solidified medium surface and incubated at 37 °C for 12–15 h. Results were calculated by evaluating the zone of inhibition in millimeters for each compound and compared with sparaxin, used as a standard (Pervez et al., 2008).

Urease Inhibition

Investigation of the urease inhibition activity of compounds 4a–e was carried according to the procedure outlined by Khan (2017) with slight modifications. Briefly, a total 200 μL (0.5 mM) reaction mixture containing a 25-μL urease enzyme solution was incubated for 15 min at 30 °C. Then, 55 μL of buffer and 100 mM of substrate urea were incubated at the same conditions. After that, 0.5% w/v sodium hydroxide, 0.1 % sodium hypochlorite (70 μL), 1 % w/v phenol, and 0.005 % w/v sodium nitroprusside (45 μL) reagent were added and incubated at 30 °C for 50 min. To conclude the urease inhibition activity, ammonia production was measured by the indophenol method. Optical density (OD) measurements were conducted at 630 nm with the aid of an ELISA plate reader (KHAN, 2017). In this assay, thiourea was used as a standard and the percentage of inhibition was calculated using the following formula:

\[
\text{% inhibition} = 100 + \frac{OD_{test}}{OD_{control}} \times 100
\]

Results And Discussion

Chemistry

To accomplish this task, the solid support acid catalyst has been prepared with silica gel and sulfuric acid. Si-H₂SO₄ catalyst was confirmed by the strong absorption band at 1060 cm⁻¹, assigned to the transverse optic (TO) and longitudinal optic (LO) modes of Si-O-Si asymmetric stretching vibrations in the FTIR spectra as depicted in Fig. 1S and 2S (images available in supplementary material) (Musić, Filipović-Vinceković, & Sekovanić, 2011). The surface morphology and texture of Si-H₂SO₄ catalyst was examined by scanning electron microscopic (SEM) images. Pure silica shows uniform crystalline form having same morphology throughout the texture whereas the Si-H₂SO₄ catalyst displayed inhomogeneity order of SiO₂ crystals due to hydrogen bonding, and appeared with amorphous topology (supplementary material Fig. 3S and Fig. 4S). The Si-H₂SO₄ catalyst was successfully employed for the synthesis of five novels 2,4,6-triarylpyridine derivatives using various aryl-aldehydes (R₁), and results are shown in Table 1.

Table 1. Si-H₂SO₄ catalyzed synthesis of 2,4,6-triarylpyridine derivatives 4a–e
Proposed Mechanism involved in the synthesis of compounds 4a–e

The proposed mechanism for the construction of 2,4,6-triarylpyridine ring involved aldol condensation of 2-acetyl furan enol (I) with 2-acetyl thiophene to furnish α,β-unsaturated ketone (III) catalyzed by Si-H₂SO₄. Then, activated thiophene enamine (V) was added to the double bond of (III) in a Micheal addition fashion to form (VI), which ultimately converts to its keto form (VII). This intermediate then undergoes heteroannulation by a nucleophilic attack of nitrogen on the carbonyl carbon to afford triaryl dihydropyridine (VIII), which upon oxidation yields the desired products (Fig. 1).

Optimization of Reaction Conditions

In order to find the optimum loading of catalyst, equimolar amounts of 2-acetyl furan, 2-acetyl thiophene, benzaldehyde, and NH₄OAc (1.5 mole %) were mixed. In this regard, three parameters were explored for reaction optimization including catalyst ratio, catalyst amount, and temperature. Initially, several attempts were made with various ratios of Si-H₂SO₄ catalyst (Table 2). Results showed that the Si-H₂SO₄ catalyst with 1:0.5 ratio proves an efficient catalyst, which led to a single spot on TLC for 2,4,6-triarylpyridines synthesis with minimum time as compared to other ratios.

Table 2. Comparison of different Si-H₂SO₄ catalyst performance in model reaction

| Entry | Ar. | Products | Time (h) | Yield % | M.P °C |
|-------|-----|----------|----------|---------|--------|
| 1     |     | 4a       | 1        | 60      | 113    |
| 2     |     | 4b       | 1        | 65      | 112    |
| 3     |     | 4c       | 1        | 63      | 110    |
| 4     |     | 4d       | 1        | 64      | 145    |
| 5     |     | 4e       | 1        | 55      | 120    |
Then, the effect of catalyst amount has been scrutinized, which revealed that with increased amount, reaction rate and product yields also increased. Results indicated that 1 g of the catalyst was sufficient for the target compound. However, 1.2 g decreased the product formation, which could be due to increased acidity of reaction system. Finally, the effect of temperature was investigated by carrying out the same model reaction at different temperatures, where good yield obtained at 80 °C (Table 3).

**Table 3. Effect of Temperature on product yield**

| T (°C) | Time (h) | % yield |
|--------|----------|---------|
| 70     | 4        | 30      |
| 75     | 3        | 45      |
| 80     | 1        | 60      |
| 85     | 1        | 40      |
| 90     | 1        | 30      |

Finally, reusability of the catalyst is one of the most significant benefits of this strategy that makes the method cost effective. In this respect, the catalyst was filtered off, washed with water, dried, and reused for the same reaction. The catalyst could be recycled at least four times with only slight reduction in the catalyst activity.

**Biological Activities**

**In Vitro Cytotoxicity**

The synthesized derivatives, 4a–e, were evaluated for cytotoxicity assay against 3T3 (mouse fibroblast) cell lines and exhibited IC<sub>50</sub> > 30, suggesting they are non-cytotoxic. Experimental results were compared with the standard drug cyclohexamide with an IC<sub>50</sub> value of 0.19 ± 0.11 μM.

**Antibacterial Screening**
We evaluated the antibacterial activity of compounds $4a$–$e$ against four Gram-negative ($Escherichia coli$, $Pseudomonas aeruginosa$, $Salmonella typhi$, $Shigella flexneri$) and two Gram-positive ($Bacillus subtilis$, $Staphylococcus aureus$) bacterial strains. Results revealed that compounds $4a$, $4c$, $4d$, and $4e$ exhibit significant antibacterial activity against Gram-positive pathogens as shown in Table 4.

| Compounds | Gram-negative bacteria (% inhibition) | Gram-positive (% inhibition) |
|-----------|-------------------------------------|-------------------------------|
|           | $E. coli$ | $P. aeruginosa$ | $S. typhi$ | $S. flexneri$ | $B. subtilis$ | $S. aureus$ |
| $4a$      |   _       |   _               |   _       |   _   | 78  | 68   |
| $4b$      |   _       |   _               |   _       |   _   |   _ |   _   |
| $4c$      |   _       |   _               | 9         |   _   | 5   | 46   |
| $4d$      |   _       | 8                | 8         |   _   | 10  | 45   |
| $4e$      |   _       |   _               | 3         |   _   | 8   | 62   |
| Sparaxin  | 92        | 93               | 93        |   _   | 94  | 90   |

Urease Inhibition Activity

Listed in Table 5 are the urease inhibitory activity of compounds $4a$–$e$. Results reveal that compound $4a$ exhibits potent urease inhibition with an IC$_{50}$ value of $12.8 \pm 1.04 \mu M$ when compared with the standard inhibitor, thiourea ($IC_{50} = 21.0 \pm 0.23 \mu M$). Results also show that compound $4a$ exhibits remarkable inhibition activity, and is more potent than the standard drug. This could be due to $\pi$-$\pi$ stacking interactions of the phenyl group at C-4 of the pyridine ring. Inhibition activity of compound $4b$ is lower than that of $4a$ with an IC$_{50}$ of $19.6 \pm 1.68 \mu M$; this might be due to the presence of an electron-donating group (–OH ) at the ortho position of phenyl ring bonded to C-4. Similarly, compound $4c$ exerted significan inhibition with an IC$_{50}$ of $15.8 \pm 2.23 \mu M$, due to the methoxy group at the para position of phenyl moiety attached to C-4 of the pyridine ring. On the other hand, the inhibition activity of compound $4d$ could be ascribed to the presence of hydrophilic interactions of furan and thiophene moieties. Furthermore, $\pi$-$\pi$ stacking interactions of the substituents may lead to the significant activity of compound $4d$, which has an IC$_{50}$ value of $17.9 \pm 1.31 \mu M$ as compared to standard. Finally, compound $4e$ was the least active as urease inhibitor with an IC$_{50}$ of $23.7 \pm 0.23 \mu M$; this may be due to decreased $\pi$-$\pi$ stacking interactions due to phenyl and chloro moieties.

Table 5. In-vitro urease inhibitory activity of triaryl pyridine derivatives
| Compound | IUPAC Name                                                                 | % Inhibition | IC$_{50}$ ± SEM [$\mu$M] |
|----------|----------------------------------------------------------------------------|--------------|--------------------------|
| 4a       | 2-(furan-2-yl)-4-phenyl-6-(thiophen-2-yl)pyridine                         | 87.9         | 12.8 ± 1.04              |
| 4b       | 4-(2-(furan-2-yl)-6-(thiophen-2-yl)pyridine-4-yl)phenol                   | 90.3         | 19.6 ± 1.68              |
| 4c       | 2-(furan-2-yl)-4-(4-methoxy phenyl)-6-(thiophen-2-yl)pyridine             | 88.9         | 15.8 ± 2.23              |
| 4d       | 2,4-di(furan-2-yl)-6-9(thiophen-2-yl)pyridine                             | 93.6         | 17.9 ± 1.31              |
| 4e       | 4-(4-chlorophenyl)-2-(furan-2-yl)-6-(thiophen-2-yl)pyridine               | 86.3         | 23.7 ± 0.23              |
| Standard | Thiourea                                                                  | 98.4         | 21.0 ± 0.23              |

**Electronic Properties**

The frontier molecular orbitals (FMOs) including the highest occupied molecular orbitals (HOMOs and HOMOs-1) and lowest unoccupied molecular orbitals (LUMOs and LUMOs+1) of compounds 4a–e at the B3LYP/TZ2P level are shown in Fig. 2. In 4a and 4b, the spatial distribution of HOMO-1 is on thiophene and furan units along with the lone pair of electrons on the nitrogen atom of pyridine, HOMO and LUMO+1 at the 2-(furan-2-yl)-6-(thiophen-2-yl)pyridine moiety while the LUMO at 4-phenyl-6-(thiophen-2-yl)pyridine. The intramolecular charge-transfer (ICT) was observed from HOMO-1 → LUMO as well as HOMO-1 → LUMO (furan to phenyl ring). In this respect, substitution of −OH at the o-position of phenyl (4b) has no significant effect on the FMOs charge density distribution, which is comparable to that of the parent compound 4a. However, introduction of −OCH$_3$ at the p-position of the phenyl group in 4c caused a significant effect on the FMOs, suggesting that HOMO-1 is delocalized at the methoxyphenyl-pyridine moiety, HOMO at 2-(furan-2-yl)-6-(thiophen-2-yl) pyridine, whereas the LUMO and LUMO+1 are localized at the thiophen-pyridine and furan-pyridine moieties, respectively.

In 4c, ICT was perceived from HOMO-1 → LUMO (methoxyphenyl to thiophene), HOMO-1 → LUMO+1 (methoxyphenyl to furan-pyridine), HOMO → LUMO (furan to thiophen-pyridine) and HOMO → LUMO+1 (thiophen to furan-pyridine). On the other hand, introduction of a furan ring to replace the phenyl group in 4d affected the spatial distribution of FMOs as compared to 4a. The HOMO-1 is delocalized on 4-(furan-2-yl)pyridine, HOMO at 2-(furan-2-yl)-6-(thiophen-2-yl)pyridin while LUMO is localized at 4-(furan-2-yl)pyridine and LUMO+1 at 2-(furan-2-yl)-6-(thiophen-2-yl)pyridine. In 4d, ICT was perceived from HOMO-1 → LUMO+1 (4-furan to thiophene and 2-furan), and HOMO → LUMO (thiophene and 2-furan to 4-furan). Substitution of the electron-withdrawing group −Cl at phenyl $para$ position of parent compound furan ring in place of the phenyl in compound 4e has no significant effect on the spatial distribution of FMOs as compared to 4a. The charge density distribution and ICT in 4e are almost similar to those of the parent compound.
Energies of FMOs, i.e., HOMO-1 (E_{HOMO-1}), HOMO (E_{HOMO}), LUMO (E_{LUMO}), LUMO+1 (E_{LUMO+1}), and HOMO-LUMO energy gaps (E_{gap}) are important parameters in the electronic and photo-physical properties of organic compounds. The E_{HOMO-1}, E_{HOMO}, E_{LUMO}, E_{LUMO+1}, and E_{gap} of compounds 4a–e at the B3LYP/TZ2P level at S_0 are listed in Table 7. Data show that E_{HOMO-1} at S_0 decreases in the order: 4d (–6.22) > 4b (–6.27) = 4c (–6.27) > 4a (–6.45). Similarly, data reveal that E_{HOMO} follows the order: 4b (–5.74) > 4e (–5.84) = 4a (–5.85) > 4d (–5.85) > 4c (–5.91), whereas the trend in the E_{LUMO} is: 4b (–1.55) > 4a (–1.67) > 4c (–1.69) > 4d (–1.78) > 4e (–1.82). In the meantime, E_{LUMO+1} decreases in the following order: 4b (–1.39) > 4d (–1.51) > 4a (–1.52) > 4c (–1.57) > 4e (–1.64), whereas E_{gap} follows the order: 4c (4.22) > 4b (4.19) = 4a (4.18) > 4d (4.07) > 4e (4.02).

Shown in Table 7 are results pertaining to FMOs energies of compounds 4a–e. Results show that substituting the electron donating groups (EDG) –OH or OCH\_3 at the para position of phenyl increases E_{HOMO-1}. Similarly, replacing the 4-phenyl with 4-furan moiety in 4d increases E_{HOMO-1}. On the other hand, the EWG –Cl at the para position of the phenyl group decreases E_{HOMO-1} as compared to 4a, whereas The EDG –OH/OCH\_3 in 4b increases E_{HOMO} as compared to the parent compound. Moreover, the EDG –OH in 4b increases E_{LUMO} whereas –OCH\_3, –Cl, and 4-furan lower that term as compared to 4a. In a similar fashion, the –OH in 4b increases E_{LUMO+1} while –OCH\_3, and –Cl lower the values as compared to 4a. Finally, the EDG –OCH\_3 causes an increase in E_{gap} while 4-furan and –Cl reduce its value as compared to compound 4a.

**Table 7.** The ground state energies of FMOs; HOMO-1 (E_{HOMO-1}), HOMO (E_{HOMO}), LUMO (E_{LUMO}), LUMO+1 (E_{LUMO+1}), and HOMO–LUMO energy gaps (E_{gap}) (in eV) of studied compounds.

| Compounds | E_{HOMO-1} | E_{HOMO} | E_{LUMO} | E_{LUMO+1} | E_{gap} |
|-----------|------------|----------|----------|------------|--------|
| 4a        | –6.37      | –5.85    | –1.67    | –1.52      | 4.18   |
| 4b        | –6.27      | –5.74    | –1.55    | –1.39      | 4.19   |
| 4c        | –6.27      | –5.91    | –1.69    | –1.57      | 4.22   |
| 4d        | –6.22      | –5.85    | –1.78    | –1.51      | 4.07   |
| 4e        | –6.45      | –5.84    | –1.82    | –1.64      | 4.02   |

**Molecular Electrostatic Potential**

The Molecular electrostatic potential (MEP) explores the reactivity of various compounds and is measured experimentally by diffraction methods (Politzer & Truhlar, 2013; Stewart, 1979) [23, 24] as well as by computational means. MEP illustrates the wide-ranging electronic and nuclear charge distribution,
which is an appropriate feature to understand the reactivity of various species (Murray & Politzer, 2011). Depicted in Fig. 3 is the MEP mapping of compounds 4a–e, illustrated in color visualizations. The red color denotes higher negative potential regions, which are required for electrophilic attack, whereas the blue color identifies the higher positive potential regions favorable for nucleophilic attack. In this respect, the MEP decreases in the order blue > green > yellow > orange > red; the red color shows the strongest repulsion while blue elucidates sufficient attraction.

In 4a, the negative electrostatic potential is located on the oxygen of furan and nitrogen of pyridine, whereas the positive electrostatic potential is found on the phenyl and thiophene moieties. In 4b, substituting the –OH at the ortho position of phenyl enhances the negative electrostatic potential at the entire molecule. Similarly, substituting the –OCH₃ at the para position leads to negative electrostatic potential on oxygen of furan, nitrogen of pyridine, and oxygen of –OCH₃, and positive electrostatic potential at phenyl and thiophene moieties in 4c. On the other hand, the positive or negative electrostatic potentials in 4d and 4e are almost similar. These results reveal that in the case of a nucleophilic attack, significant repulsion would be at oxygen and nitrogen atoms in the furan and pyridine moieties, and on oxygen of –OCH₃ in all studied compounds, except for 4b where most of the sites would behave as repulsion. The significant attraction would be thiophene and phenyl rings. It is also expected that in the case of an electrophilic attack, significant repulsion would be at the thiophene and phenyl rings whereas the noteworthy attraction would be at oxygen and nitrogen atoms in furan and pyridine, and on –Cl and oxygen of –OCH₃.

Hirshfeld Charges

Shown in Fig. 3 are the Hirshfeld charges of the newly synthesized compounds 4a–e. Hirshfeld charges on various carbon, oxygen, nitrogen, and hydrogen atoms in the 2-(furan-2-yl)-6-(thiophen-2-yl)pyridine moiety are almost similar in all of these derivatives. However, substituting the –OH or –OCH₃ increases the positive charge of the carbon atom. The Hirshfeld charge on C2 was found –0.043 in 4a while introducing the electron donating group (EDG) –OH lead to change the charge to 0.081 in 4b. Similarly, Hirshfeld charge on C4 was –0.044 in 4a while substituting with the EDG –OCH₃ lead to changing the value to 0.080. The Hirshfeld charge on C3 was found –0.044 in 4a while by introducing EDG –OH lead to change its charge –0.073 in 4b. In addition, Hirshfeld charge on C3 was –0.044 in 4a while by substituting EDG –OCH₃ lead to change its value, i.e., –0.062. The Hirshfeld charge on C5 was found –0.044 in 4a while by introducing EDG –OH/OCH₃ lead to change its charge –0.060/–0.071 in 4b / 4c, which is due to the donating effect of –OH/OCH₃. Replacing 4-phenyl by 4-furan moiety reduced the positive charge at C7 from 0.015 to 0.007 in 4c. The electron-withdrawing group (EWG) –Cl at the para position of the phenyl group reduced the negative charge at C2, C3, C4, C5 and C6 atoms. However, the most significant effect was observed at C4 where the negative charge –0.044 has become 0.018 due to the withdrawing effect of –Cl. The Hirshfeld charges revealed that –OH and –OCH₃ at the ortho and para positions increase the negative charges on C3 and C5 atoms, suggesting that more negative charge on these sites in 4b and 4c would be favorable for electrophilic substitution.
Conclusions

In summary, we have synthesized five new 2,4,6-triarylpyridine derivatives in good yield using the solid support catalyst approach. The present methodology offered attractive features such as short reaction time, simplicity, and reusability of the catalyst. The prepared compounds were tested in vitro for antibacterial and urease inhibition activities. Results revealed that all of these newly synthesized compounds display good to moderate antibacterial and urease inhibition potential, and may serve as lead molecule in the drug discovery of new antibacterial drugs. In addition, the electronic properties of these derivatives have been determined with various methods such as DFT and Hirshfeld charges method.

List Of Abbreviations

\[\text{[SiO}_2\text{-VO(OH)}_2\]| = Heterogeneous silica vanadic acid  \\
\text{[TS-1]} = \text{Titanium silicates}  \\
\text{[MTT]} = 3\text{-[4, 5-dimethylthiazole-2-yl]-2, 5-diphenyl-tetrazolium bromide}  \\
\text{[FBS]} = \text{Fetal bovine serum}  \\
\text{[EDG]} = \text{Electron donating group}  \\

Declarations

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Table
The Table 6 is not available.

**Scheme**

Scheme 1 is available in the Supplementary Files section.

**Figures**

**Figure 1**

Proposed Mechanism for a one-pot Synthesis of 2,4,6-triarylpyridines
Figure 2

Ground state charge density of some frontier molecular orbitals (FMOs) of compounds 4a–e (contour value = 0.035)
Figure 3

Molecular electrostatic potential surfaces views of triarylpypyridines 4a-e.
Figure 4

Hirshfeld charges of compounds 4a-e

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.
- Scheme1.png
- Supplymentrymaterial1.docx