An efficient synthesis and antimicrobial evaluation of some new pyrazoline, pyrimidine and benzodiazepine derivatives bearing 1,3,5-triazine core

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ABSTRACT. In our present investigation a new class of diverse sets of acetyl pyrazolines (6a-e), amino pyrimidines (7a-e) and 1,5-benzodiazepines (8a-e) bearing 1,3,5-triazine core were synthesised from chalcones (5a-e). Treatment of chalcone with hydrazine hydrate, guanidine hydrochloride and o-phenylenediamine afforded the corresponding acetyl pyrazoline, amino pyrimidine and 1,5-benzodiazepine derivatives respectively. The structures of all the newly synthesised compounds were assigned on the basis of FTIR, $^1$H NMR, $^{13}$C NMR, mass spectral data as well as elemental analysis. In vitro antimicrobial proficiency of the title compounds were assessed against selected pathogens S. aureus MTCC 96, S. pyogenes MTCC 442, E. coli MTCC 443 and P. aeruginosa MTCC 1688 bacteria for antibacterial activities as well as antifungal activities against C. albicans MTCC 227, A. niger MTCC 282 and A. clavatus MTCC 1323 were used. The minimum inhibitory concentration (MIC) was determined by broth dilution method and recorded at the lowest concentration inhibiting growth of the organism. Among the synthesised compounds 6b, 6c, 7b, 8b, 8d and 8e exhibited excellent antimicrobial activity and said to be the most proficient members of the series.

1. INTRODUCTION

Varieties of infectious diseases caused by microbes are increasing rapid in recent year and still a major threat to public health in the world [1], even though incredible development in medicinal chemistry. Owing to non-availability of requisite medicines and emergence of prevalent drug resistance, the impact is more delicate in developing countries. Scientists and doctors are still struggling to find solutions with various forms of medications. Thus the demand for designing of new antimicrobial agents having excellent potential with high beneficial index with novel structural classes and mechanisms is necessary. 1,3,5-Triazine has received an enormous attention among the researchers and chemists as a consequence of essential role in novel drug discovery owing to three-fold symmetry, allows for versatile modifications, uncomplicated by region chemical concerns and has proven a useful biological target with diverse pharmacological properties of the molecules [2-3]. And therefore, it is an ongoing effort to synthesise novel antimicrobial agents bearing 1,3,5-triazine core. Chalcones are compounds of chalconoid group chemically known as 1,3-diarylprop-2-en-1-one and key precursors in the synthesis of a large array of biologically important heterocycles [4-8]. It is therefore, not surprising that many synthetic methods have been developed for the preparation of heterocycles starting from chalcones precursors. Pyrazoline is an important five member heterocyclic compound containing nitrogen as a hetero atom and various methods have been worked out for their synthesis. Due to the interesting activity of variously substituted pyrazolines as biological agents considerable attention has been focused on this class of compounds. They are used as anti-inflammatory [9], antitumor [10], anticancer [11], antimicrobial [12] etc…. Among the existing various pyrazoline type derivatives, 1-acetyl pyrazolines have been identified as one of the most promising derivatives. In view of the importance of these derivatives, the title compound (6a-e) is achieved. Pyrimidine is a six member heterocyclic compound...
containing four carbon and two nitrogen atoms and is pharmacologically inactive but its synthetic derivatives possess an important role in modern medicine. In medicinal chemistry pyrimidine derivatives have been very eminent for their therapeutic applications like anti-infective [13], antibacterial [14], analgesic [15], diuretic [16] etc... 1,5-Benzodiazepine is a vital nitrogen containing seven member heterocyclic compound in pharmaceutical research because they possess a variety of bioactivities. Various methods have been worked out for their synthesis and numerous derivatives have been published in the literature with a diverse range of biological activities such as antimicrobial [17], anticonvulsant, antianxiety, analgesic, sedative [18] as well as cholecystokinin - 2 receptor antagonists [19]. In light of the above considerations, we herein report the reaction of 2-(3'-trifluromethylphenylamino)-4-(tetrahydro-1',4'-oxazine)-6-[4'-(3'-aryl-2''-propenon-1''-yl)phenylamino]-1,3,5- triazines (5a-e) with hydrazine hydrate, guanidine hydrochloride and o-phenylenediamine furnished subsequent conversion to products acetyl pyrazolines (6a-e), amino pyrimidines (7a-e) and 1,5-benzodiazepines (8a-e) derivatives respectively.

2. EXPERIMENTAL

2. 1. Material

All the chemicals and solvents used for reaction were of analytical reagent (AR) grade. All the melting points were resolute in open capillary method and are uncorrected. IR spectra were recorded on Shimadzu FTIR 8401 spectrophotometer using potassium bromide pellets. $^1$H NMR and $^{13}$C NMR spectra were recorded on a Bruker Avance 400 F (MHz) spectrometer (Bruker Scientific Corporation Ltd., Switzerland) using CDCl$_3$, DMSO as a solvent and TMS as an internal standard at 400 MHz. Chemical shifts are reported in parts per million (ppm) and coupling constant (J) are reported in Hertz. Elemental analysis was carried out by Perkin-Elmer 2400 series-II elemental analyser (Perkin-Elmer, USA). Mass spectra were scanned on a Shimadzu LCMS 2010 spectrometer (Shimadzu, Tokyo, Japan). Purity of the compounds were checked by thin layer chromatography using TLC aluminum sheets Silica Gel 60 F 254 (Merck) plates of 0.25 mm thickness and detection of the components were made by exposure to UV light or keeping the plates in iodine chamber.

2. 2. Method

2. 2 A. General procedure for the compounds (A), (B), (C) and chalcones (5a-e)

The starting precursors A, B and C were prepared by the reported procedure [20]. The chalcones (5a-e) were prepared by the reported method [12] in good yields by a base catalysed Claisen-Schmidt condensation of appropriately substituted benzaldehydes and substituted ketone (C).

2. 2 B. Preparation of 2 - (3' - trifluromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {1'' - acetyl - 5'' - (4'' - methoxyphenyl) - 2'' - pyrazolin 3'' - yl} phenylamino] - 1,3,5 - triazines (6a)

A 100 ml round bottomed flask, fitted with a reflux condenser was charged with a mixture of chalcone (5a) (0.01 mol, 5.76 gm in 30 ml ethanol) and hydrazine hydrate (0.015 mol, 0.75 gm in 5 ml ethanol). To make the mixture acidic catalytic amount of glacial acetic acid (5 ml) was added. The reaction mixture was heated under reflux temperature for 5-6 hours. The progress of the reaction was investigated by TLC using toluene: methanol (12:6 v/v) eluent as mobile phase. After completion of the reaction, the mixture was cooled to room temperature then poured into crushed ice and neutralised with Na$_2$CO$_3$. The solid mass separated was collected by filtration, washed well with hot water and recrystallised from methanol to get product (BI) in good yield with high purity. Similarly, the remaining compounds (6b-e) were synthesised by this given method.
2. 2 C. 2 - (3'- trifluromethylphenylamino) - 4 - (tetrahydro - 1', 4'- oxazine) - 6 - [4' - {2'' - amino - 6'' - (4'' - methoxyphenyl) - pyrimidin - 4'' - yl} phenylamino] - 1,3,5 - triazines (7a)

Compound (5a) (0.01 mol, 5.76 gm in 30 ml ethanol) and guanidine hydrochloride (0.015 mol, 1.43 gm in 5 ml ethanol) dissolved in ethanol was mixed in 100 ml round bottomed flask. To make this mixture alkaline 40% KOH (5 ml) was added to the reaction mixture and refluxed for 4-5 hours. The progress of the reaction was investigated by TLC using toluene: methanol (15:9 v/v) as mobile phase. In the same way, the remaining compounds (7b-e) were prepared by this method. All the newly synthesised compounds (7a-e) were characterised by IR, 1H NMR, 13C NMR and mass spectroscopy as well as elemental analysis. The characteristic data of the entire synthesised compounds are given in spectral and physical analysis data (3).

2. 2 D. Preparation of 2 - (3'- trifluromethylphenylamino) - 4 - (tetrahydro - 1', 4'- oxazine) - 6 - [4' - {4'' - (4'' - methoxyphenyl) - 3' H - benzo - 1'' 5'' - diazepin - 2'' - yl} phenylamino] - 1,3,5 - triazines (8a)

Compound (5a) (0.01 mol, 5.76 gm in 30 ml ethanol) condensed with o- phenylenediamine (0.01 mol, 1.0 gm) in the presence of catalytic amount of glacial acetic acid (5 ml) in ethanol at refluxed temperature for 5-6 hours in 100 ml round bottomed flask. The progress of the reaction was monitored by TLC using toluene: methanol (12:8 v/v) eluent as mobile phase. After completion of the reaction, the reaction mixture was poured into crushed ice and neutralised with dilute HCl. Finally, the product was filtered, washed with water, dried and recrystallised from methanol to get product (7a) in good yield with high purity. Similarly, the remaining compounds (7b-e) were prepared by this method.

In the same way, the remaining compounds (8a-e) were prepared by this method.

All the newly synthesised compounds (6a-e), (7a-e) and (8a-e) were characterised by IR, 1H NMR, 13C NMR and mass spectroscopy as well as elemental analysis. The spectral and physical analysis data of the entire synthesised compounds are given in spectral analysis data (3).

2. 3. Reaction Scheme

Methodical synthetic route for the target compounds (6a-e), (7a-e) and (8a-e)
3. SPECTRAL ANALYSIS DATA

**Compound (6a):** 2 - (3' - trifluromethylphenylamino) - 4 - (tetrahydro - 1',4' - oxazine) - 6 - [4' - {1'' - acetyl 5'' - (4'' - methoxyphenyl)2'' - pyrazolin 3'' - yl} phenylamino] - 1,3,5- triazine: Yield: 78%; mp: 147 °C; IR (KBr, νmax, cm⁻¹): 3358 (-NH str.), 3007 (=CH str. aromatic), 2900 (C-H str. pyrazoline moiety), 1610 (C=O str.), 1583 (C=N str. pyrazoline moiety), 1546 (-NH bend.), 1510 (C=C str. aromatic), 1365 (CH3 str.), 1247 (asymmetric C-O-C str. ether linkage), 1176 (OCH3 str.), 1070 (C-F str.), 833 (C-H bend. 1.4 substituted benzene ring), 800 (C=N str. 1,3,5- triazine); 1H NMR (400MHz, CDCl3, δ ppm): 2.4 (s, 3H, -COCH3), 3.0 (dd, J = 9.7 & 15.4 Hz, 1H, -CH2-CH), 3.6 (dd, J = 9.7 & 12.4 Hz, 1H, -CH2-CH), 5.6 (dd, J = 4.0 & 11.5 Hz, 1H, -CH2-CH2-Ar), 3.94 (s, 3H, 4-OCH3), 3.73 (concealed t, 4H, -CH2, oxazine ring), 3.84 (concealed t, 4H, -CH2, oxazine ring), 6.7 - 8.1 (m, 13H, 12 Ar-H and 1- NH); 13C NMR (400 MHz, DMSO, δ ppm): 23.4 (CH3), 40.9 (CH2, methylene, pyrazoline moiety), 48.9 (CH2, oxazine), 55.9 (OCH3), 65.4 (CH-Ar), 66.9 (CH2, oxazine), 111.4 (CH), 115.1 (CH), 116.7 (CH), 121.3 (CH), 123.6 (CF3), 126.5 (CH), 126.9 (C), 127.7 (CH), 128.5 (CH), 129.7 (CH), 130.2 (CH), 131.8 (C), 141.7 (C), 142.2 (C), 142.8 (C), 146.2 (C), 151.6 (C=N of pyrazoline moiety), 166.2 (C=N of 1,3,5- triazine), 168.7 (C=N of 1,3,5-triazine), 168.5 (C=O), 175.4 (C=N of 1,3,5- triazine); M.S. (m/z): 632.5 (M⁺); Anal. Calcd. for C33H31N8F3O3: C, 64.76; H, 4.93; N, 17.71%. Found: C, 60.79; H, 4.90; N, 17.69%.

**Compound (6b):** 2 - (3' - trifluromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {1'' - acetyl 5'' - (3'' - chlorophenyl) - 2'' - pyrazolin 3'' - yl} phenylamino] - 1,3,5- triazine: Yield: 71%; mp: 129 °C; IR (KBr, νmax, cm⁻¹): 3289 (-NH str.), 3078 (=CH str. aromatic), 2956 (C-H str. pyrazoline moiety), 1678 (C=O str.), 1597 (C=N str. pyrazoline moiety), 1551 (-NH bend.), 1489 (C=C str. aromatic), 1370 (CH3 str.), 1220 (asymmetric C-O-C str. ether linkage), 1089 (C-F str.), 826 (C-H bend. of 1,4 substituted benzene ring), 809 (C=N str. 1,3,5- triazine), 659 (C-Cl); 1H NMR (400MHz, CDCl3, δ ppm): 2.1 (s, 3H, -COCH3), 3.3 (dd, J = 11.2 & 16.5 Hz, 1H, -CH2-CH), 3.9 (dd, J = 12.3 & 16.5 Hz, 1H, -CH2-CH), 5.1 (dd, J = 4.5 & 12.4 Hz, 1H, -CH2-CH2-Ar), 3.53 (concealed t, 4H, -CH2, oxazine ring), 3.78 (concealed t, 4H, -CH2, oxazine ring), 7.1 - 8.3 (m, 13H, 12 Ar-H and 1- NH); 13C NMR (400 MHz, DMSO, δ ppm): 24.7 (CH3), 39.4 (CH2, methylene, pyrazoline moiety), 46.1 (CH2, oxazine), 64.2 (CH-Ar), 65.8 (CH2, oxazine), 110.8 (CH), 114.7 (CH), 115.2 (CH), 122.2 (CH), 124.7 (CF3), 125.1 (CH), 126.3 (C), 128.4 (CH), 129.1 (CH), 130.5 (CH), 131.4 (CH), 132.0 (C), 134.3 (C), 140.9 (C), 141.5 (C), 143.2 (C), 150.3 (C=N of pyrazoline moiety), 162.1 (C=N of 1,3,5- triazine), 166.5 (C=N of 1,3,5- triazine), 167.9 (C=O), 172.0 (C=N of 1,3,5- triazine); M.S. (m/z): 637.9 (M⁺); Anal. Calcd. for C33H28N8F3O2Cl: C, 58.45; H, 4.43; N, 17.59%. Found: C, 58.44; H, 4.41; N, 17.58%.

**Compound (6c):** 2 - (3' - trifluromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {1'' - acetyl 5'' - (3'' - phenoxyphenyl) - 2'' - pyrazolin 3'' - yl} phenylamino] - 1,3,5- triazine: Yield: 79%; mp: 139 °C; IR (KBr, νmax, cm⁻¹): 3156 (-NH str.), 3016 (=CH str. aromatic), 2947 (C-H str. pyrazoline moiety), 1656 (C=O str.), 1590 (-NH bend.), 1578 (C=N str. pyrazoline moiety), 1460 (C=C str. aromatic), 1367 (CH3 str.), 1268 (C-O-C str. phenoxy), 1254 (asymmetric C-O-C str. ether linkage), 1099 (C-F str.), 836 (C-H bend. of 1,4 substituted benzene ring), 804 (C=N str. 1,3,5- triazine); 1H NMR (400MHz, CDCl3, δ ppm): 2.9 (s, 3H, -COCH3), 3.0 (dd, J = 10.8 & 15.2 Hz, 1H, -CH2-CH), 3.4 (dd, J = 9.6 & 15.2 Hz, 1H, -CH2-CH), 4.8 (concealed d, 1H, -CH2-CH2-Ar), 3.67 (concealed t, 4H, -CH2, oxazine ring), 3.92 (concealed t, 4H, -CH2, oxazine ring), 6.7 - 8.4 (m, 18H, 17 Ar-H and 1- NH); 13C NMR (400 MHz, DMSO, δ ppm): 23.9 (CH3) 39.7 (CH2, methylene, pyrazoline moiety), 48.1 (CH2, oxazine), 66.2 (CH-Ar), 67.9 (CH2, oxazine), 110.6 (CH), 112.4 (CH), 114.1 (CH), 116.5 (CH), 119.1 (CH), 121.2 (CH), 123.7 (CF3), 125.4 (CH), 126.8 (C), 127.1 (CH), 129.2 (CH), 129.9 (CH), 131.6 (CH), 132.4 (CH), 133.0 (C), 137.7 (C), 141.8 (C), 142.1 (C), 148.9 (C=N of pyrazoline moiety), 156.7 (C), 157.3 (C), 163.2 (C=N of 1,3,5- triazine), 166.8 (C=N of 1,3,5- triazine), 169.1 (C=O), 173.5 (C=N of 1,3,5- triazine); M.S. (m/z): 694.3 (M⁺); Anal. Calcd. for C37H33N8F3O3: C, 63.97; H, 4.78; N, 16.13%. Found: C, 63.95; H, 4.75; N, 16.16%.
Compound (6d): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {1''- acetyl 5'' - (2'' - nitrophenyl) 2'' - pyrazolin 3''- yl} phenylamino] - 1,3,5 - triazine; Yield: 69%; mp: 119 ¹C; IR (KBr, ν max, cm⁻¹): 3316 (ν-NH str), 3058 (=CH str. aromatic), 2919 (C-H str. pyrazoline moiety), 1637 (C=O str.), 1597 (ν-NH bound.), 1590 (C=N, str. pyrazoline moiety), 1539 (C-NO₂), 1503 (C=C str. aromatic), 1369 (CH₃ str.), 1221 (asymmetric C-O-C str. ether linkage), 1026 (C-F str.), 831 (C-H bend. 1.4 disubstituted benzene ring), 799 (C-N str. 1.3,5- triazine); ¹H NMR (400MHz, CDCl₃, δ ppm): 2.6 (s, 3H, -COCH₃), 3.3 (dd, J = 9.3 & 15.9 Hz, 1H, -CH₂-CH), 3.9 (dd, J = 9.3 & 13.4 Hz, 1H, -CH₂-CH₂-), 4.2 (dd, J = 4.5 & 12.2 Hz, 1H, -CH-CH₂-Ar), 3.51 (concealed t, 4H, -CH₂, oxazine ring), 3.73 (concealed t, 4H, -CH₂, oxazine ring), 7.0 - 8.2 (m, 13H, 12 Ar-H and 1-NE); ¹³C NMR (400 MHz, DMSO, δ ppm): 23.9 (CH₃), 41.3 (CH₂, methylene, pyrazoline moiety), 48.6 (CH₂, oxazine), 65.6 (CH-Ar), 67.1 (CH₂, oxazine), 111.2 (CH), 113.8 (CH), 114.5 (CH), 120.3 (CH), 122.8 (CF₃), 124.9 (CH), 125.7 (C), 127.3 (CH), 128.1 (CH), 129.0 (CH), 131.8 (CH), 133.7 (C), 137.6 (C), 140.9 (C), 142.7 (C), 148.3 (C), 152.1 (C=N of pyrazoline moiety), 163.4 (C=N of 1.3,5- triazine), 167.6 (C=N of 1.3,5- triazine), 169.2 (C=O), 173.8 (C=N of 1.3,5- triazine); M.S. (m/z): 647.4 (M⁺); Anal. Calcd. for C₃₁H₂₈N₈F₆O₃: C, 57.50; H, 4.35; N, 19.47%. Found: C, 57.53; H, 4.39; N, 19.44%.

Compound (6e): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {1''- acetyl 5'' - phenyl 2'' - pyrazolin 3''- yl} phenylamino] - 1,3,5 - triazine; Yield: 70%; mp: 156 ¹C; IR (KBr, ν max, cm⁻¹): 3187 (ν-NH str), 3008 (=CH str. aromatic), 2987 (C-H str. pyrazoline moiety), 1648 (C=O str.), 1591 (ν-NH bound.), 1583 (C=N str. pyrazoline moiety), 1497 (C=C str. aromatic), 1372 (CH₂ str.), 1239 (asymmetric C-O-C str. ether linkage), 1064 (C-F str.), 838 (C-H bend. 1.4 disubstituted benzene ring), 803 (C=N str. 1.3,5- triazine); ¹H NMR (400MHz, CDCl₃, δ ppm): 1.9 (s, 3H, -COCH₃), 2.6 (dd, J = 11.8 & 16.2 Hz, 1H, -CH₂-CH₂-), 3.1 (concealed d, 1H, -CH₂-CH₂-), 3.7 (dd, J = 4.7 & 11.6 Hz, 1H, -CH-CH₂-Ar), 3.39 (concealed t, 4H, -CH₂, oxazine ring), 3.61 (concealed t, 4H, -CH₂, oxazine ring), 7.2 - 8.3 (m, 14H, 13 Ar-H and 1-NE); ¹³C NMR (400 MHz, DMSO, δ ppm): 25.7 (CH₃), 38.2 (CH₂, methylene, pyrazoline moiety), 46.5 (CH₂, oxazine), 64.6 (CH-Ar), 66.8 (CH₂, oxazine), 109.7 (CH), 111.2 (CH), 113.7 (CH), 117.4 (CH), 124.1 (CF₃), 124.9 (CH), 126.2 (C), 127.8 (CH), 129.3 (CH), 130.7 (CH), 132.3 (CH), 134.4 (C), 138.6 (C), 141.2 (C), 142.0 (C), 149.6 (C=N, pyrazoline moiety), 162.4 (C=N of 1.3,5- triazine), 165.9 (C=N of 1.3,5- triazine), 170.8 (C=O), 172.6 (C=N of 1.3,5- triazine); M.S. (m/z): 602.6 (M⁺); Anal. Calcd. for C₃₁H₂₀N₈F₆O₂: C, 61.79; H, 4.85; N, 18.6%. Found: C, 61.82; H, 4.82; N, 18.62%.

Compound (7a): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {2''- amino - 6'' - (4'' - methoxypenyl) - pyrimidin - 4''- yl} phenylamino] - 1,3,5 - triazine; Yield: 73%; mp: 143 ¹C; IR (KBr, ν max, cm⁻¹): 3307 (=NH str. 1° amine of pyrimidine moiety), 3109 (=NH str.), 3064 (=CH str. aromatic), 1672 (C=N str. pyrimidine moiety), 1577 (=NH bound.), 1510 (C-N str. aromatic), 1247 (asymmetric C-O-C str. ether linkage), 1165 (OCH₃ str.), 1068 (C-F str.), 831 (C-H bend. 1.4 disubstituted benzene ring), 802 (C-N str. of 1.3,5- triazine); ¹H NMR (400MHz, CDCl₃, δ ppm): 3.89 (s, 3H, 4-OCH₃), 3.43 (concealed t, 4H, -CH₂, oxazine ring), 3.69 (concealed t, 4H, -CH₂, oxazine ring), 5.18 (s, 2H, -NH₂), 6.9 - 8.2 (m, 14H, 13 Ar-H and 1-NH); ¹³C NMR (400 MHz, DMSO, δ ppm): 47.3 (CH₂ oxazine), 55.6 (OCH₃) 68.7 (CH₂ oxazine), 101.7 (CH, pyrimidine moiety), 114.8 (CH), 115.1 (CH), 116.4 (CH), 116.8 (CH), 121.1 (CH), 123.8 (CF₃), 125.8 (C), 128.1 (C), 128.3 (CH), 128.8 (CH), 129.3 (CH), 131.4 (C), 138.6 (C), 142.2 (C), 160.6 (C), 161.1 (C=N of 1.3,5- triazine), 162.0 (C, pyrimidine moiety), 163.8 (C=N of pyrimidine moiety), 165.2 (C of pyrimidine moiety), 166.4 (C=N of 1.3,5- triazine), 172.6 (C=N of 1.3,5- triazine); M.S. (m/z): 615.1 (M⁺); Anal. Calcd. for C₃₃H₂₆N₈F₆O₃: C, 60.48; H, 4.58; N, 20.48%. Found: C, 60.52; H, 4.56; N, 20.51%.

Compound (7b): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - {2''- amino - 6'' - (3'' - chlorophenyl) - pyrimidin - 4''- yl} phenylamino] - 1,3,5 - triazine; Yield: 70%; mp: 130 ¹C; IR (KBr, ν max, cm⁻¹): 3390 (=NH str. 1° amine of pyrimidine moiety), 3209 (=NH str. 1° amine of pyrimidine moiety).
Compound (7c): 2 - (3' - trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - 2'' - amino - 6'' - (3'' - phenoxy) - pyrimidin - 4'' - yl] phenylamino] - 1,3,5 - triazine; Yield: 69%; mp: 165 °C; IR (KBr, v max, cm⁻¹): 3368 (-NH str. of pyrimidine moiety), 3226 (-NH str.), 3034 (=CH str.); 1H NMR (400MHz, CDCl₃, δ ppm): 3.79 (concealed t, 4H, -CH₂, oxazine ring), 3.97 (concealed t, 4H, -CH₂, oxazine ring), 4.9 (s, 2H, -NH₂), 6.8 - 8.3 (m, 19H, 18 Ar-H and 1 -NH); 13C NMR (400 MHz, DMSO, δ ppm): 48.3 (CH₂ oxazine), 65.9 (CH₂ oxazine), 99.1 (CH, pyrimidine moiety), 113.4 (CH), 114.8 (CH), 115.3 (CH), 116.7 (CH), 118.7 (CH), 120.1 (CH), 121.8 (CH), 123.5 (CF₃), 126.2 (C), 127.0 (C), 128.1 (CH), 128.6 (CH), 130.2 (CH), 131.3 (CH), 133.0 (C), 138.4 (C), 140.3 (C), 157.6 (C), 157.9 (C), 164.6 (C=N of 1,3,5- triazine), 165.3 (C=N of pyrimidine moiety), 167.2 (C, pyrimidine moiety), 169.0 (C, pyrimidine moiety), 172.2 (C=N of 1,3,5- triazine), 173.5 (C=N of 1,3,5- triazine); M.S. (m/z): 677.0 (M⁺); Anal. Calcd. for C₃₀H₂₅N₇F₃O₂Cl: C, 58.11; H, 4.03; N, 20.30%.

Compound (7e): 2 - (3' - trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4' - oxazine) - 6 - [4' - 2'' - amino - 6'' - (3'' - phenoxazine) - pyrimidin - 4'' - yl] phenylamino] - 1,3,5 - triazine; Yield: 64%; mp: 125 °C; IR (KBr, v max, cm⁻¹): 3326 (-NH str. of pyrimidine moiety), 3229 (-NH str.), 3036 (=CH str. aromatic), 1629 (C=N str. pyrimidine moiety), 1549 (-NH bend.), 1472 (C=C str. aromatic), 1251 (asymmetric C-O-C str. ether linkage), 1091 (C-F str.), 843 (C-H bend. 1.4 disubstituted benzene ring), 829 (C-H bend. 1.4 disubstituted benzene ring), 806 (C=N str. 1,3,5- triazine), 1H NMR (400MHz, CDCl₃, δ ppm): 3.33 (concealed t, 4H, -CH₂, oxazine ring), 3.68 (concealed t, 4H, -CH₂, oxazine ring), 5.4 (s, 2H, -NH₂), 6.6 - 8.0 (m, 14H, 13 Ar-H and 1 -NH); 13C NMR (400 MHz, DMSO, δ ppm): 47.6 (CH₂ oxazine), 64.2 (CH₂ oxazine), 102.9 (CH, pyrimidine moiety), 113.1 (CH), 114.5 (CH), 115.7 (CH), 116.8 (CH), 119.3 (CH), 122.8 (CF₃), 125.4 (C), 126.6 (C), 127.7 (CH), 128.9 (CH), 131.2 (CH), 132.5 (C), 142.4 (C), 143.8 (C), 146.0 (C), 162.7 (C=N of pyrimidine moiety), 164.9 (C=N of 1,3,5- triazine), 165.0 (C, pyrimidine moiety), 168.2 (C, pyrimidine moiety), 168.8 (C=N of 1,3,5- triazine), 174.2 (C=N of 1,3,5- triazine); M.S. (m/z): 630.8 (M⁺); Anal. Calcd. for C₃₀H₂₅N₇F₃O₂C: 63.81; H, 4.46; N, 18.60%. Found: C, 63.79; H, 4.51; N, 18.57%.
Compound (8a): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4'- oxazine) - 6 - [4'-
{4''- (4'''- methoxyphenyl) - 3'' H - benzo - 1''', 5'' diazepin - 2''- yl] phenylamino} - 1,3,5 - triazine:
Yield: 70%; mp: 107 °C; IR (KBr, v max, cm⁻¹): 3346 (-NH str.), 3067 (=CH str. aromatic), 1610
(C=N str. benzodiazepine moiety), 1581 (-NH bend.), 1494 (C=C str. aromatic), 1241 (asymmetric
C-O-C str. ether linkage), 1157 (OCH3 str.), 1100 (C-F str.), 848 (C-H bend. 1.4 disubstituted
benzene ring), 805 (C=N str. 1,3,5- triazine); 1H NMR (400MHz, CDCl3, δ ppm): 2.72 (dd, J = 6.2 &
13.0 Hz, 1H, CH3, diazepine moiety), 2.44 (dd, J = 6.2 & 14.2 Hz, 1H, CH3, diazepine moiety),
3.78 (s, 3H, 4-OCH3), 3.44 (concealed t, 4H, -CH2, oxazine ring), 3.74 (concealed t, 4H, -CH2,
oxazine ring), 6.9 - 8.1 (m, 17H, 16 Ar-H and 1 - NH); 13C NMR (400 MHz, DMSO, δ ppm): 37.8
(CH3, methylene, diazepine moiety), 49.8 (CH2, oxazine), 57.9 (4 - OCH3), 69.7 (CH2, oxazine),
110.4 (CH), 113.8 (CH), 116.3 (CH), 118.7 (CH), 120.1 (CH), 122.5 (CH), 124.4 (CF3), 127.3(C),
128.5 (CH), 130.2 (CH), 131.8 (C), 132.1 (CH), 133.1 (CH), 137.8 (C), 141.3 (C), 143.8 (C), 149.2
(C), 157.3 (C=N), 162.7 (C), 166.2 (C=N of 1,3,5- triazine), 169.4 (C=N of 1,3,5- triazine), 171.4
(C=N of 1,3,5 triazine); M.S. (m/z): 664.0 (M+); Anal. Calcd. for C35H31N8F3O2: C, 65.05; H,
4.70; N, 16.86%. Found: C, 65.01; H, 4.74; N, 16.83%.

Compound (8b): 2 - (3'- trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4'- oxazine) - 6 - [4'-
{4''- (3'''- chlorophenyl) - 3'' H - benzo - 1''', 5'' diazepin - 2''- yl] phenylamino} - 1,3,5 - triazine:
Yield: 70%; mp: 107 °C; IR (KBr, v max, cm⁻¹): 3378 (-NH str.), 3034 (=CH str. aromatic), 1599
(C=N str. benzodiazepine moiety), 1583 (-NH bend.), 1496 (C=C str. aromatic), 1249 (asymmetric
C-O-C str. ether linkage), 1039 (C-F str.), 835 (C-H bend. 1.4 disubstituted benzene ring), 801
(C=N str. 1,3,5- triazine), 650 (C=Cl); 1H NMR (400MHz, CDCl3, δ ppm): 2.87 (concealed d, 1H,
CH3, diazepine moiety), 3.26 (concealed d, 1H, CH3, diazepine moiety), 3.56 (concealed t, 4H, -CH2,
oxazine ring), 3.83 (concealed t, 4H, -CH2, oxazine ring), 7.2 - 8.3 (m, 17H, 16 Ar-H and 1 - NH);
13C NMR (100 MHz, DMSO, δ ppm): 38.2 (CH3, methylene, diazepine moiety), 48.6 (CH2, oxazine),
67.3 (CH2, oxazine), 111.6 (CH), 112.5 (CH), 115.1 (CH), 116.0 (CH), 118.3 (CH), 120.2
(CH), 123.7 (CF3), 126.2 (C), 129.1 (CH), 131.6 (CH), 132.4 (C), 133.9 (CH), 134.7 (CH), 135.0
(C), 138.2 (C), 142.8 (C), 144.1 (C), 147.9 (C), 155.8 (C=N of diazepine moiety), 164.1 (C=N of
1,3,5- triazine), 167.0 (C=N of 1,3,5- triazine), 172.9 (C=N of 1,3,5- triazine); M.S. (m/z): 669.4
(M+); Anal. Calcd. for C36H33N8F3O2: C, 65.05; H, 4.70; N, 16.86%. Found: C, 65.01; H, 4.74; N,
16.83%.

Compound (8c): 2 - (3' - trifluoromethylphenylamino) - 4 - (tetrahydro - 1', 4'- oxazine) - 6 - [4'-
{4''- (3'''- phenoxyphenyl) - 3'' H - benzo - 1''', 5'' diazepin - 2''- yl] phenylamino} - 1,3,5 - triazine:
Yield: 80%; mp: 126 0°C; IR (KBr, v max, cm⁻¹): 3362 (-NH str.), 3059 (=CH str. aromatic), 1594
(C=N str. benzodiazepine moiety), 1587 (-NH bend.), 1480 (C=C str. aromatic), 1291 (C-O-C str.
phenoxy), 1226 (asymmetric C-O-C str. ether linkage), 1049 (C-F str.), 842 (C-H bend. 1.4 disubstituted
benzene ring), 798 (C=N str. 1,3,5- triazine); 1H NMR (400MHz, CDCl3, δ ppm): 3.45
(dd, J = 6.4 & 13.5 Hz, 1H, CH3, diazepine moiety), 3.78 (dd, J = 7.0 & 13.5 Hz, 1H, CH3,
diazepine moiety), 3.36 (concealed t, 4H, -CH2, oxazine ring), 3.56 (concealed t, 4H, -CH2, oxazine ring),
6.8 - 8.2 (m, 22H, 21 Ar-H and 1 - NH); 13C NMR (400 MHz, DMSO, δ ppm): 39.8 (CH2,
methylene, diazepine moiety), 46.1 (CH2, oxazine), 66.9 (CH2, oxazine), 109.1 (CH), 111.3 (CH),
113.8 (CH), 115.7 (CH), 117.0 (CH), 118.2 (CH), 119.6 (CH), 121.4 (CH), 124.2 (CF3), 125.6 (C),
127.1 (CH), 128.0 (CH), 130.9 (CH), 131.3 (C), 132.4 (CH), 133.8 (CH), 136.0 (C), 140.2 (C),
142.6 (C), 145.9 (C), 153.6 (C=N of diazepine moiety), 156.3 (C), 158.4 (C), 165.6 (C=N of 1,3,5-
triazine), 166.3 (C=N of 1,3,5- triazine), 170.2 (C=N of 1,3,5- triazine); M.S. (m/z): 726.6 (M+);
Anal. Calcd. for C41H33N8F3O2: C, 67.76; H, 4.57; N, 15.42%. Found: C, 67.74; H, 4.53; N,
15.45%.
Compound (8d): 2 - (3’-trifluoromethylphenylamino) - 4 - (tetrahydro - 1’, 4’- oxazine) - 6 - [4’- {4”- (2”- nitrophenyl) - 3” H - benzo - 1”, 5”- diazipen - 2”- yl} phenylamino] - 1,3,5 - triazine: Yield: 65%; mp: 117 °C; IR (KBr, νmax, cm⁻¹): 3306 (-NH str.), 3071 (=CH str. aromatic), 1592 (C=N str. benzodiazepine moiety), 1577 (-NH bend.), 1539 (C-NO₂), 1502 (C=C str. aromatic), 1219 (asymmetric C-O-C str. ether linkage), 1085 (C-F str.), 831 (C-H bend. 1.4 substituted benzene ring), 806 (C=N str. 1,3,5- triazine); ¹H NMR (400MHz, CDCl₃, δ ppm): 2.96 (concealed d, 1H, CHₓ, diazepine moiety), 3.54 (dd, J = 6.8 & 14.2 Hz, 1H, CHₓ, diazepine moiety), 3.78 (concealed t, 4H, -CH₂, oxazine ring), 4.02 (concealed t, 4H, -CH₂, oxazine ring), 7.0 - 8.1 (m, 17H, 16 Ar-H and 1 - NH); ¹³C NMR (400 MHz, DMSO, δ ppm): 38.6 (CHₓ, methylene, diazepine moiety), 48.3 (CH₂, oxazine), 68.6 (CH₂, oxazine), 110.4 (CH), 112.1 (CH), 114.7 (CH), 115.8 (CH), 117.2 (CH), 119.8 (CH), 124.8 (CF₃), 125.9 (C), 127.4 (CH), 130.0 (CH), 133.7 (C), 134.2 (CH), 135.1 (CH), 137.9 (C), 140.9 (C), 143.2 (C), 145.6 (C), 150.9 (C), 156.5 (C=N of diazepine moiety), 165.8 (C=N of 1,3,5- triazine), 168.8 (C=N of 1,3,5- triazine), 174.1 (C=N of 1,3,5-triazine); M.S. (m/z): 679.0 (M⁺); Anal. Calcd. for C₅₃H₄₉NₓF₃O₅: C, 61.83; H, 4.19; N, 18.55%. Found: C, 61.83; H, 4.19; N, 18.52%.

Compound (8e): 2 - (3’-trifluoromethylphenylamino) - 4 - (tetrahydro - 1’, 4’- oxazine) - 6 - [4’- {4”- (phenyl) - 3” H - benzo - 1”, 5”- diazipen - 2”- yl} phenylamino] - 1,3,5 - triazine: Yield: 72%; mp: 100 °C; IR (KBr, νmax, cm⁻¹): 3319 (-NH str.), 3061 (=CH str. aromatic), 1598 (C=N str. benzodiazepine moiety), 1571 (-NH bend.), 1489 (C=C str. aromatic), 1243 (asymmetric C-O-C str. ether linkage), 1054 (C-F str.), 844 (C-H bend. 1.4 disubstituted benzene ring), 803 (C=N str. 1,3,5-triazine); ¹H NMR (400MHz, CDCl₃, δ ppm): 3.64 (dd, J = 7.2 & 14.2 Hz, 1H, CHₓ, diazepine moiety), 4.03 (dd, J = 7.2 & 13.9 Hz, 1H, CHₓ, diazepine moiety), 3.49 (concealed t, 4H, -CH₂, oxazine ring), 3.81 (concealed t, 4H, -CH₂, oxazine ring), 6.8 - 8.3 (m, 18H, 17 Ar-H and 1 - NH); ¹³C NMR (400 MHz, DMSO, δ ppm): 40.9 (CHₓ, methylene, diazepine moiety), 46.2 (CHₓ, oxazine), 66.1 (CH₂, oxazine), 108.1 (CH), 110.7 (CH), 112.2 (CH), 113.4 (CH), 115.6 (CH), 117.1 (CH), 122.3 (CF₃), 126.7 (C), 128.9 (CH), 130.8 (CH), 131.6 (CH), 133.4 (C), 135.2 (CH), 136.0 (CH), 138.8 (C), 142.1 (C), 144.4 (C), 148.9 (C), 154.0 (C=N of diazepine moiety), 162.1 (C=N of 1,3,5- triazine), 166.4 (C=N of 1,3,5- triazine), 170.2 (C-N, 1,3,5 - triazine); M.S. (m/z): 634.7 (M⁺); Anal. Calcd. for C₃₅H₂₉NₓF₃O₅: C, 66.24; H, 4.60; N, 17.66%. Found: C, 66.27; H, 4.58; N, 17.63%.

4. RESULTS AND DISCUSSION

4.1. Chemistry

The aim of the present study was to develop an efficient protocol with good to excellent yields in a short span of time without formation of any side product. The formation of designed compounds was confirmed by their IR, ¹H NMR and ¹³C NMR spectral and elemental analysis. The IR spectrum of compound 6a exhibited a strong band at 1610 cm⁻¹ for C=O stretching of acetyl group attached at N₁ position in pyrazoline ring. The C–H stretching was observed at 2900 cm⁻¹. Characteristic band of C=N stretching of pyrazoline moiety appeared at 1583 cm⁻¹. A strong absorption band was observed at 1365 cm⁻¹ due to the presence of the CH₃ group. The aromatic C-H bending vibrations for 1,4 disubstituted benzene ring and C=N stretching of 1,3,5-triazine core were observed at 833 and 800 cm⁻¹ respectively. The ¹H NMR spectrum of compound 6a showed a singlet at δ 2.4 ppm for the COCH₃ protons. The pro-chiral methylene protons Cₓ”-H of pyrazoline appeared as two distinct doubles of a doublet at δ 3.0 ppm (J = 9.7 & 15.4 Hz) and at δ 3.6 ppm (J = 9.7 & 12.4 Hz) for the CHₓ”-CH and CHₓ’-CH protons, thereby indicating that both the protons are magnetically non-equivalent and diastereotopic. The chiral Cₓ”-H proton of pyrazoline appeared as a doubles of a doublet at δ 5.6 ppm (J = 4.0 & 11.5 Hz) due to CH-CH₂-Ar proton. The other remaining twelve aromatic protons appeared as a multiplet signal at δ 6.7–7.0 ppm. In the ¹³C NMR spectrum of compound 6a, the signal appeared at δ 23.4 ppm was assigned to the methyl carbon and the most deshielded signal that appeared at δ 168.5 ppm was assigned to the carbonyl carbon of the acetyl group attached with the pyrazoline unit. The signals for aromatic carbons appeared between δ
111.4-146.2 ppm in the $^{13}$C spectrum. The IR spectrum of compound 7a showed a strong characteristic band at 1672 cm$^{-1}$ and 3307 cm$^{-1}$ due to the C=N and NH$_2$ group of pyrimidine ring. The aromatic C=C stretching and C-H bending vibrations for 1,4 disubstituted benzene ring were appeared at 1510 and 831 cm$^{-1}$ respectively. The $^1$H NMR spectrum of compound 7a showed a singlet at δ 5.18 ppm due to NH$_2$ proton on pyrimidine nucleus which confirmed the cyclisation of pyrimidine moiety. The other remaining thirteen aromatic protons resonate as a multiplet signal at δ 6.9-8.0 ppm. $^{13}$C NMR spectrum of compound 7a showed a signal at δ 163.8 ppm assigned to the C=N carbon of pyrimidine ring which assigned the pyrimidine unit. The signals for aromatic carbons appeared between δ 114.8-142.6 ppm in the $^{13}$C spectrum. The IR spectrum of compound 8a exhibited a strong band at 1556 cm$^{-1}$ due to the C=N of diazepine unit. Aromatic C=C and C-H bending vibrations for 1,4 disubstituted benzene ring were appeared at 1494 and 848 cm$^{-1}$ respectively. The $^1$H NMR spectrum of compound 8a showed C$_3^\equiv$H as doublet of doublet at δ 2.72 ppm (J = 6.2 & 13.0 Hz) and at δ 3.44 ppm (J = 6.2 & 14.2 Hz). The other remaining seventeen aromatic protons resonate as a multiplet signal at δ 6.9-8.0 ppm. In the $^{13}$C NMR spectrum of compound 8a showed a shielded signal at δ 37.8 signal for methylene carbon and the most deshielded signal appeared at 157.3 ppm due to the C=N of diazepine ring which confirmed diazepine moiety. The signals for aromatic carbons appeared between δ 110.4-149.2 ppm in the $^{13}$C spectrum. There are no absorptions in the region of 1600 -1700 cm$^{-1}$ in IR spectra of compound 7a and 8a which indicating the absence of a C=O group of chalcone moiety in these structures and further confirmed the cyclisation of chalcone in to its derivatives. Moreover, distinctive singlet around δ 3.65 - 4.21 ppm stands for methoxy group of aryl ring attached to pyrazoline, pyrimidine and diazepine unit, singlet around δ 8.1 - 8.5 ppm stands for secondary amine attached with 1,3,5-triazine. Further, mass spectra of compound 6a, 7a and 8a showed molecular ion peak M+ at m/z 632.5, 615.1 and 664.0 (100%) along with other fragment peaks, which also supported the structure of compound 6a, 7a and 8a respectively. The obtained elemental analysis values within ±0.4% of theoretical values of all the synthesised compounds were good agreement with the molecular formula.

4.2. Antimicrobial evaluation
The antibacterial and antifungal activity of newly synthesised compounds (6a-e), (7a-e) and (8a-e) was carried out by broth dilution method [21] according to National Committee for Clinical Laboratory Standards (NCCLS, 2002). It is one of the non automated in vitro bacterial / fungal susceptibility tests. This classic method yields a quantitative result for the amount of antimicrobial agents that is needed to inhibit growth of specific microorganisms which is carried out in tubes. Upon reviewing antimicrobial data (Table 1) it has been observed that in Gram positive bacterial strains, compounds 7b (MIC = 50 µg/mL), 8d and 8e (MIC = 62.5 µg/mL) exhibited excellent activity against Staphylococcus aureus (MTCC 96) compared to Ampicillin (MIC = 250 µg/mL) and equipotent activity to Chloramphenicol and Ciprofloxacin (MIC = 50 µg/mL) while 8c (MIC = 100 µg/mL), 6a, 6d, 6e, 7c and 7d (MIC = 125 µg/mL) were found to be more potent compared to Ampicillin (MIC = 250 µg/mL) and less potent to Chloramphenicol and Ciprofloxacin (MIC = 50 µg/mL) against Staphylococcus aureus (MTCC 96). Compounds 6b, 6c and 8b (MIC = 62.5 µg/mL) were found to possess excellent activity compared to Ampicillin (MIC = 250 µg/mL) and equivalent to Chloramphenicol and Ciprofloxacin (MIC = 50 µg/mL) against Streptococcus pyogenes (MTCC 442) whereas compounds 6a, 6d, 6e, 7b and 8e (MIC = 100 µg/mL), 7c, 7d and 8d (MIC = 125 µg/mL) were found equally potent to Ampicillin (MIC = 100 µg/mL) and less potent to Chloramphenicol and Ciprofloxacin (MIC = 50 µg/mL) against Streptococcus pyogenes (MTCC 442). Compounds 6b, 7a and 8a (MIC = 200 µg/mL) were found to be more active than Ampicillin (MIC = 250 µg/ml) while compounds 6c, 7e and 8b (MIC = 250 µg/mL) were exerted equally potent to Ampicillin (MIC = 250 µg/mL) and modest to Chloramphenicol and Ciprofloxacin (MIC = 50 µg/mL) against Staphylococcus aureus. On the other hand in Gram negative bacterial strains, compound 6b (MIC = 50 µg/mL) showed excellent potency against Escherichia coli (MTCC 443) compared to Ampicillin (MIC = 100 µg/mL), comparable to Chloramphenicol (MIC = 50 µg/mL) and modest to Ciprofloxacin (MIC = 25 µg/mL) whereas
compounds 6e, 7c, 8b and 8e (MIC = 100 µg/mL), 6c, 8a and 8d (MIC = 125 µg/mL) were found to have comparable potency to Ampicillin (MIC = 100 µg/mL) against *Escherichia coli* (MTCC 443). Compounds 6b and 8e (MIC = 100 µg/mL), 7c and 8b (MIC = 125 µg/mL) exerted equally active to Ampicillin (MIC = 100 µg/mL) and less active to Chloramphenicol (MIC = 50 µg/mL) and Ciprofloxacin (MIC = 25 µg/mL) against *Pseudomonas aeruginosa* (MTCC 441). The antifungal screening data (Table 1) revealed that compound 8b(MIC = 100 µg/mL) exhibited excellent activity against *Candida albicans* (MTCC 227) compared to Greseofulvin (MIC = 500 µg/mL) and significant to Nystatin (MIC = 100 µg/mL) whereas compounds 6c, 6d, 7a, 7b, 7d, 8a, 8c and 8d (MIC = 500 µg/mL) were found to have equipotent activity to Greseofulvin (MIC = 500 µg/mL) against *Candida albicans* (MTCC 227). None of the compound showed promising antifungal activity against *Aspergillus niger* (MTCC 282) and *Aspergillus clavatus* (MTCC 1323).

Table 1. In Vitro antimicrobial activity of synthesised compounds (6a-e), (7a-e) and (8a-e)

<table>
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<th>Minimal fungicidal concentration MIC (µg/mL)</th>
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<td>S. p MTCC 442</td>
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</table>

Sa.: *Staphylococcus aureus*; Sp.: *Streptococcus pyogenes*; Ec.: *Escherichia coli*; Pa.: *Pseudomonas aeruginosa*; Ca.: *Candida albicans*; An.: *Aspergillus niger*; Ac.: *Aspergillus clavatus*; A: Ampicillin; B: Chloramphenicol; C: Ciprofloxacin; D: Greseofulvin; E: Nystatin. MTCC: Microbial Type Culture Collection. ‘-’ represents ‘not tested’.
5. CONCLUSION

In summary, the synthesis of biologically important three elegant protocols (acetyl pyrazolines, amino pyrimidines and 1,5-benzodiazepines) has been developed using the potential of chalcones with the hope of generating new pharmacologically important molecules that could be useful as potent antimicrobial agents. The method reported in this investigation is effective in giving excellent conversion to the product, less-energy consuming and apparently substituent insensitive. There is no necessary to do any extra or special work during isolation of product. The results indicated that all the derivatives of new chemical entities exhibited appreciable activities against multidrug-resistant bacteria. Among the fifteen newly synthesised compounds, analogs 6b, 6c, 7b, 8b, 8d and 8e were found to be the most proficient members of the series. It is interesting to note that compounds bearing the electron withdrawing chloro group in the aryl moiety 6b, 7b and 8b showed the highest antibacterial and antifungal activities. Further in depth SAR study of the active scaffolds and the mechanism research are also required. These result suggest that chalcones and their derivatives have an opportunity to behave as generation of newer antimicrobial agents and have excellent scope for further development as commercial antimicrobial agents.

Acknowledgement

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References


