Synthesis and antimicrobial activity of imidazo- and pyrimido[2,1-f]-theophyllines

Mosselhi A. N. Mosselhi¹, Elham S. Darwish¹, Klaus Peseke²

¹ Department of Chemistry, Faculty of Science, Cairo University, Cairo, Egypt

² Department of Chemistry, Faculty of Science, Rostock University, Rostock, Germany

Received 12 July 2007; Accepted 12 November 2007; Published online 11 April 2008 © Springer-Verlag 2008

Abstract Heating of 8-aminotheophylline with methyl (Z)-2-benzoylamino-3-(dimethylamino)propenoate in acetic acid afforded in a one-pot synthesis a new pyrimido[2,1-f]theophylline derivative. Methylation of this by using CH₃I/NaH furnished in good yield the double methylated derivative. Furthermore, glycosidation of the former with $1-\alpha$ -bromo-2,3,4,6-tetra-*O*-acetyl-D-glucose gave the β -glucoside derivative. Reaction of 8-aminotheophylline with [bis(methylthio)methylene]malonitrile, ethyl[bis(methylthio)methylene]cyanoacetate, 1,3-diphenylprop-2-en-1-one, 2-cyano-1,3-diphenylprop-2-en-1-one, 1-(4-nitrophenyl)-3-(dimethylamino)prop-2-ennitrile, 1-phenyl-3-(dimethylamino)prop-2-en-1-one, 2-substituted 3-aryl or heteroarylprop-2-ennitrile and ethyl(arylmethylene)cyanoacetate in N,N-dimethylformamide in the presence of anhydrous potassium carbonate afforded also the corresponding new derivatives of pyrimido-[2,1-f]theophylline. However, 8-aminotheophylline reacted in similar manner with 3-chloropentan-2,4dione and 2-bromo-1-phenylethanone to give the corresponding imidazo[2,1-f]theophyllines. Furthermore, azo-coupling of one of these with 4-methylphenyldiazonium chloride was performed. The antimicrobial activity of the products has been evaluated. The structures of all new compounds obtained were established by their spectral analyses.

Keywords Theophylline; Fused purines; Methylxanthines; Glycoside; Antimicrobial activity.

Introduction

Among new alkylxanthines, 7- and 8-substituted derivatives were investigated in respect of their bronchospasmolytic [1–4], anticancer [5], and circulatory blood system activity [6]. A large amount of work has been performed on the fused systems derived from theophylline, including synthetic procedures and structure determination [7–15] but only few of the synthesized new heterocyclic derivatives were pharmacologically tested, which revealed antiinflammatory [16], anti P-388 leukemia [17], and vascular relaxing agents [18]. Recently, it has been found that anellation of a six or seven membered ring at the 7,8-positions of theophylline changed the profile of its CNS activity [19, 20].

In literature several examples of [*f*]-fused purines have been reported including pyrrolo[2,1-*f*] [21], oxazolo[2,3-*f*] [22, 23], imidazo[2,1-*f*] [24–26], pyrido[2,1-*f*] [21], pyrimido[2,1-*f*] [21, 27–32, 36], oxazino[2,3-*f*] [33], pyrazino[2,1-*f*] [21], diaze-pino[2,1-*f*] [20, 3], 2,4-benzodiazepino[3,2-*f*] [34], 1,2,4-triazino[3,2-*f*] [35–37], and 1,2,4-triazepino-[3,2-*f*] [35] purines. As part of our studies of new fused purine compounds as potential antimicrobial agents, we wish to report the synthesis of new derivatives of imidazo and pyrimido[2,1-*f*]purine *via*

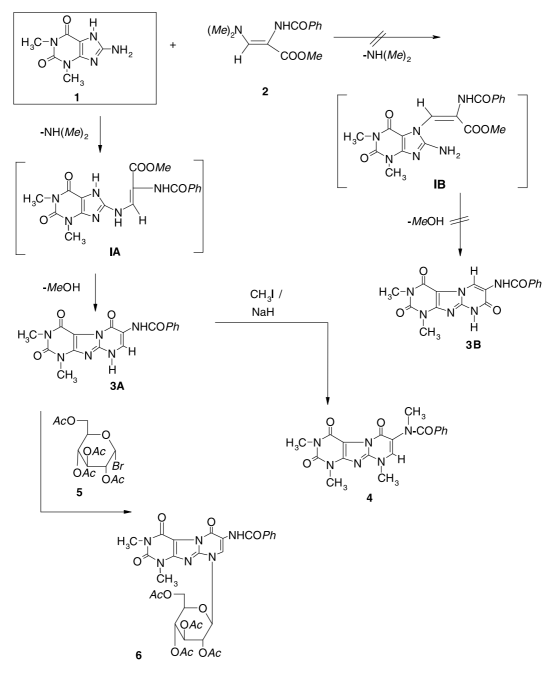
Correspondence: Mosselhi A. N. Mosselhi, Department of Chemistry, Faculty of Science, Cairo University, Cairo, Egypt. E-mail: mosselhi@hotmail.com

reaction of 8-aminotheophylline with ketene, thioacetal, propenenitrile, and haloketone derivatives.

Results and discussion

The starting 8-aminotheophylline (1) was prepared as previously reported [38]. Refluxing of 1 with methyl (Z)-2-benzoylamino-3-(dimethylamino)propenoate (2) in acetic acid for 15 h gave a single product as

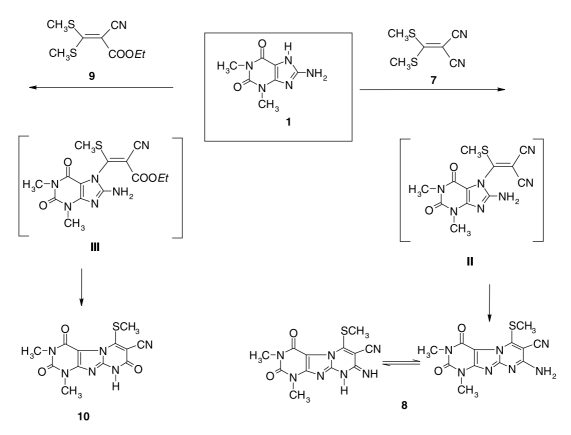
indicated by TLC analysis of the crude product. The structure of the isolated product was established on the basis of its spectral (MS, IR, and ¹H NMR) analyses. The mass spectrum of the product isolated revealed a molecular ion peak (m/z) at 366.34 of C₁₇H₁₄N₆O₄. Its infrared spectrum revealed one absorption band of NH at $\bar{\nu} = 3409 \text{ cm}^{-1}$ and no band of NH₂. Also the ¹H NMR spectrum showed 2NH signals at $\delta = 9.8$ and 13.8 ppm. Furthermore, the



Scheme 1

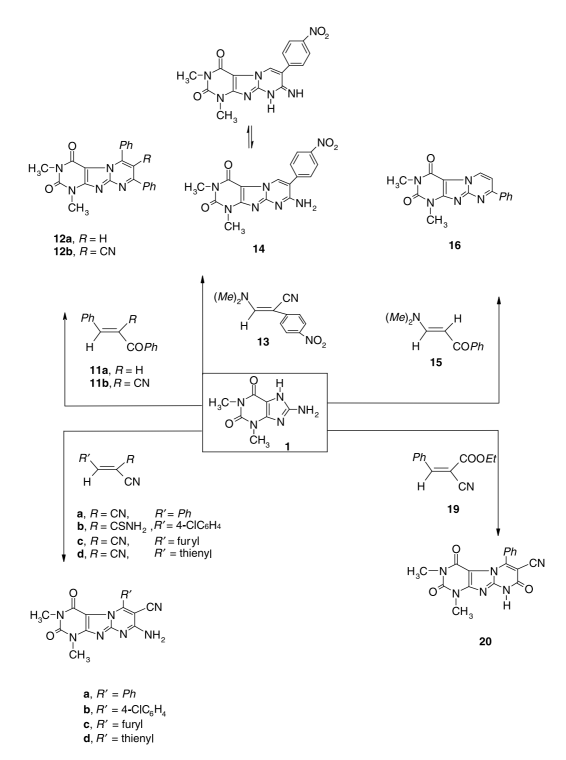
¹³C NMR spectrum of the obtained product revealed 15 carbon signals. Since the chemical shift of the carbonyl carbon at position 6 of product 3 (168.8) is similar to that of the reported carbonyl carbon at position 6 (169) of 7-benzoylamino-1,3-diphenyl-pyrido[2,3-d][1,2,4]triazolo[4,3-a]pyrimidin-5,6(1H,8H)dione [39], these spectral data were in full agreement with the expected structure of the product obtained to resemble that of pyrimido[2,1-f]theophylline derivative **3A** and not the isomer **3B** (Scheme 1). Formation of **3A** might occur *via* initial electrophilic substitution of the 8-amino group of 1 in acidic medium [39, 47], to give 1A as an intermediate which undergoes cyclization to the final product 3(Scheme 1). The assignment of the structure 3A is also substantiated by investigation of its methylation and glycosidation reactions. Thus, methylation of 3A by using methyl iodide in the presence of sodium hydride yielded the double methylation product 4. The ¹H NMR spectrum of the latter product 4 revealed two signals of new CH₃ groups at $\delta = 3.1$ (PhCON-CH₃) and 3.15 (N9-CH₃) ppm. However, glycosidation of **3** with 1- α -bromo-2,3,4,6-tetraacetyl-D-glucose (5) afforded the glucoside derivative **6** (Scheme 1). The ¹H NMR spectrum of **6** showed the anomeric proton as a doublet at $\delta = 6.2$ ppm with a spin-spin coupling constant (*J*) of 10.5 Hz corresponding to a diaxial orientation of H-1' and H-2' protons indicating the β -glucoside [45].

The reaction of 1 with a molar equivalent of 2-(bis-methylthiomethylene) malonitrile (7) and ethyl[bis(methylthio)methylene]cyanoacetate (9) in refluxing N,N-dimethylformamide (DMF) containing equivalent amounts of anhydrous potassium carbonate for 15 h (evidenced by TLC) afforded the corresponding cyclized products, pyrimido[2,1-f]theophylline derivatives 8 and 10 (Scheme 2). The structures of 8 and 10 we confirmed by spectral data. The ¹H NMR of the product 8 showed a signal for SCH_3 protons at $\delta = 2.6$ and two signals of NH at $\delta = 8.0$ and 9.6 ppm and no signal of NH₂ was observed. This finding indicates that the structure of the latter product 8 exists in imine form. The ¹H NMR of the product 10 showed one NH signal at $\delta = 9.6 \text{ ppm}$ and no signal of NH₂. Moreover, the IR spectra of 8 and 10 revealed bands characteristic for a CN group.



Scheme 2

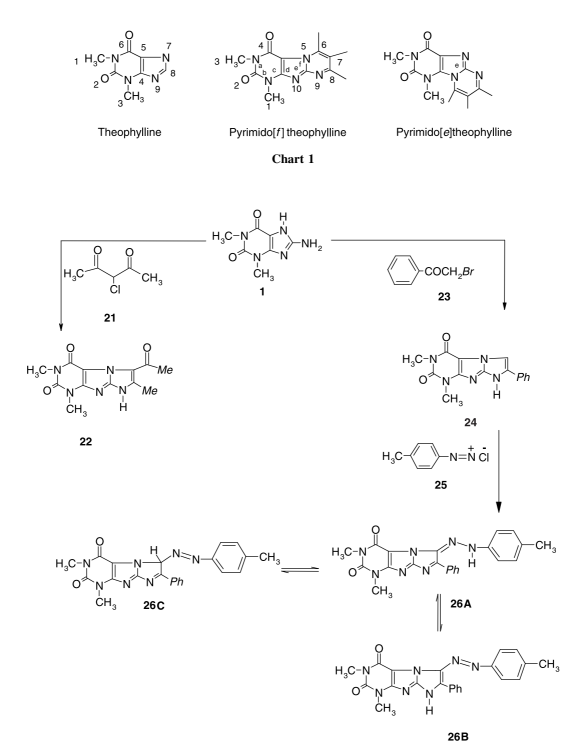
As reported in literature [21, 40], the formation of **8** and **10** may proceed *via* initial alkylation of the ring nitrogen in **1** to give **II** and **III**, respectively as intermediates which undergo cyclization to the final products **8** and **10**. The synthetic approach pointed out here was extended to enable the synthesis of other functionally substituted pyrimido-[2,1-f]theophyllines for their biological evaluation. When 8-aminotheophylline **1** was reacted with 1,3diphenylprop-2-en-1-one (**11a**), 2-cyano-1,3-diphe-



829

nylprop-2-en-1-one (11b), 1-(4-nitrophenyl)-3-dimethylaminoprop-2-ennitrile (13), and 1-phenyl-3dimethylaminoprop-2-en-1-one (15) in refluxing DMFcontaining equivalent amounts of anhydrous potassium carbonate for 10 h (TLC), the corresponding pyrimido[2,1-*f*]theophylline derivatives **12a**, **12b**, **14**, and **16** were obtained (Scheme 3).

The structures of **12**, **14**, and **16** were established on the basis of their spectral data (MS, ¹H NMR, and IR). For example, the ¹H NMR spectrum of **12a**



showed a characteristic signal at $\delta = 6.99$ ppm of the proton at position 7 and that of **14** revealed a signal at $\delta = 8.39$ ppm for the proton at position 6. Also in the ¹H NMR spectrum of product **16**, two doublet signals at $\delta = 7.54$ and 8.26 ppm with a coupling constant J = 7.5 Hz corresponding to CH=CH group of positions 6 and 7 are observed [19].

Similarly, by the reaction of 1 with 2-substituted 3-aryl or heteroarylprop-2-ennitrile 17 and ethyl-(arylmethylene)cyanoacetate 19 in refluxing DMF containing equivalent amounts of anhydrous potassium carbonate for 12h, the corresponding pyrimido[2,1-f]theophylline derivatives 18 and 20 were obtained (Scheme 3). Elemental analyses and spectral data were consistent with the proposed structures of 18 and 19. In the light of the foregoing results of all new pyrimidotheophylline derivatives 3, 8, 10, 12, 14, 16, 18, and 20 obtained, it is proposed that all isolated products are consistent with a pyrimido[f]theophylline ring system and not the other isomeric pyrimido[e]theophylline (Chart 1) [19, 21, 46]. This is due to the steric hindrance caused by the proximity of the N1-CH3 and substituents in the pyrimidine ring fused. In addition the ¹H NMR spectra of all isolated products revealed the signal of the N1–CH₃ protons at $\delta = 3.42-3.65$ ppm. This value is very close to that of N3-CH₃ of theophylline $(\delta = 3.59 \text{ ppm})$ (Chart 1).

Attempts to prepare the ring system imidazo[2,1fltheophylline were made by reacting 8-aminotheophylline (1) with 3-chloropentan-2,4-dione (21) and 2-bromo-1-phenylethanone (23) in refluxing DMF containing equivalent amounts of anhydrous potassium carbonate for 10h. The corresponding imidazo-[2,1-f]theophyllines 22 and 24 were isolated, respectively (Scheme 4). The constitutions of the products 22 and 24 were confirmed by elemental and spectral analyses. In the ¹H NMR spectrum of 24, a signal of aromatic CH at $\delta = 8.15$ ppm was observed. Treatment of 24 with 4-methylphenyl diazonium chloride (25) in ethanol containing sodium acetate at 0-5°C for 3 h afforded a single product 26 according to TLC. The structure of the latter product was elucidated by elemental analysis and spectral data. The IR spectrum revealed absorption bands of NH at $\bar{\nu} = 3451$ (NH) and two absorption bands of 2 CO at 1696 and 1643 cm⁻¹; its ¹H NMR showed a signal of NH proton at $\delta = 8.35$ ppm (D₂O exchangeable) and no signal of CH proton. Also the UV absorption spectrum of 26 in methanol revealed

two absorption bands at λ_{max} 267 and 461 nm. These findings suggest that the isolated product 26 may be a mixture of hydrazone 26A and azo tautomeric form 26B, whereas the tautomeric form 26C does not seem to play a role.

Antimicrobial activity

The compounds **3**, **4**, **8**, **10**, **12a**, **12b**, **16**, **18c**, **18d**, and **26** were evaluated for their antifungal and antibacterial activities against four fungal species namely *Aspergillus fumigatus* (*AF*), *Penicillium italicum* (*PI*), *Syncephalastrum racemosum* (*SR*), and *Candida albicans* (*CA*) as well as four bacteria species namely *Staphylococcus aureus* (*SA*), *Pseudomonas aeruginosa* (*PA*), *Bacillus subtilis* (*BS*), and *Escherichia coli* (*EC*).

The organisms were tested against the activity of solutions in a concentration of $1.0 \,\mu g/cm^3$ of each compound and using inhibition zone diameter in cm (*IZD*) as a criterion for its antimicrobial activity.

Terbinafin as an antifungal agent and chloramphenicol as an antibacterial agent were used as references to evaluate the potency of the tested compounds under the same conditions. The results are depicted in Table 1. The results revealed that some compounds, such as **12a**, **12b**, **16**, **18c**, **18d**, and **26** have no activities against the tested organisms *PA*, *BS*, and *EC*, while compounds **12a**, **16**, and **26** exhibited the highest degree of inhibition against the tested organisms *SA* and *PI*.

Table 1 Antimicrobial activity of the products 3, 4, 8, 10, 12a, 12b, 16, 18c, 18d, and 26*

Compound no.	AF	PI	SR	CA	SA	PA	BS	EC
3	0	0	0	0	+	0	+	0
4	+	0	0	+	0	0	0	0
8	0	0	0	0	+	0	+	+
10	0	0	+	0	+	0	+	0
12a	0	++	+	0	+	0	0	0
12b	0	+	0	0	+	0	0	0
16	0	+	0	0	++	0	0	0
18c	+	0	0	0	0	0	0	0
18d	0	0	0	0	+	0	0	0
26	0	++	0	+	+	0	0	0

*50 cm³ of solution in *DMF* whose concentration was $1.0 \,\mu\text{g/cm}^3$ was tested; chloramphenicol as standard antibacterial agent (*IZD* 1.0 cm); terbinfin as standard antifungal agent (*IZD* 1.0 cm); ++ *IZD* 0.6–1.0 cm; + *IZD* 0.1–0.5 cm; 0 no inhibition detected.

Experimental

IR spectra were determined on a KBr disc using a Perkin-Elmer 1650 (FT-IR) spectrophotometer, ¹H-NMR spectra were recorded on a Bruker AC 250 MHz and on a Varian Gemini 200 MHz NMR spectrometer using *TMS* as the internal reference; the mass spectra were recorded on a GC-MS spectrometer, the ionizing voltage was 70 eV. Thin layer chromatography was performed on silica gel sheets F 1550 LS 254 of Schleicher & Schüll. UV absorption spectra were recorded on a Perkin-Elmer Lambda 40 spectrophotometer. Melting points were measured on a Gallenkamp melting point apparatus. Elemental analyses were carried out at the Microanalytical Center of Cairo University and were within 0.4% of the theoretical values.

The starting materials such as 8-aminotheophylline [38] (1), methyl (Z)-2-benzoylamino-3-dimethylaminopropenoate [41] (2), [bis(methylthio)methylene]malonitrile [42] (7), ethyl[bis(methylthio)methylene]cyanoacetate [42] (9), 1,3-diphenylprop-2-en-1-one [43] (11a), 2-cyano-1,3-diphenylprop-2-en-1-one [43] (11b), 1-(4-nitrophenyl)-3-dimethyylaminoprop-2-enenitrile [44] (13), 1-phenyl-3-dimethylaminoprop-2-enenitrile [44] (15), 2-substituted 3-aryl- or -heteroarylprop-2-enenitrile [43] (17), and ethyl (arylmethylene)cyanoacetate [43] (19) were prepared by literature methods. 3-Chloropentan-2,4-dione (21) and 2-bromo-1phenylethanone (23) were bought from Aldrich.

7-Benzoylamino-1,3-dimethyl-pyrimido[2,1-f]purine-

1,2,3,4,6,9-hexahydro 2,4,6-triones (**3**, $C_{17}H_{14}N_6O_4$) A mixture of 3.90 g (0.02 mol) **1** and 4.96 g (0.02 mol) **2** in 50 cm³ glacial acetic acid was heated under reflux for 15 h. The reaction was followed by TLC using CHCl₃/CH₃OH (90/10, v/v) as eluent. The reaction solvent was evaporated *in vacuo* and the residue was recrystallized from *DMF*.

Yield 5.13 g (70%); $R_{\rm f} = 0.22$; mp > 300°C; IR: $\bar{\nu} = 3409$ (NH), 1710, 1671, 1653, 1597 (4CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.2$ (s, 3H, N3–CH₃), 3.41 (s, 3H, N1–CH₃), 7.50–8.01 (m, 5H_{arom}), 8.3 (s, 1H, =CH), 9.8 (s, 1H, NH), 13.8 (s, 1H, NH) ppm; MS: m/z (%) = 366 (M⁺, 30), 349 (5), 232 (5), 176 (3), 149 (35), 105 (100), 77 (40), 44 (25).

7-(Benzoylmethylamino)-1,3,9-trimethylpyrimido[2,1-f]-

purine-1,2,3,4,6,9-hexahydro-2,4,6-trione (**4**, $C_{19}H_{18}N_6O_4$) To a stirred suspension of 0.002 mol sodium hydride (60% oil) in 5 cm³ *DMF*, a solution of 3.7 g (0.01 mol) **3** in 10 cm³ *DMF* was added. The reaction mixture was cooled to 0–5°C and a solution of 0.0012 mol methyl iodide in 2 cm³ *DMF* was added. The reaction mixture was stirred overnight and the solvent was evaporated. The residue was treated with 10 cm³ ice-water and 2 cm³ acetic acid and then stirred 2 h. The solid product was collected, washed with water, and recrystallized from dioxane/*DMF* (1/1, v/v) to give pure colourless powder [TLC using CHCl₃/CH₃OH (90/10, v/v) as eluent].

Yield 0.3 g (75%); $R_{\rm f}$ = 0.30; mp > 300°C; IR: $\bar{\nu}$ = 1715, 1680, 1650, 1600 (4CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ = 3.1 (s, 3H, *Ph*CON–CH₃), 3.15 (s, 3H, N-9 CH₃), 3.25 (s, 3H, N3–CH₃), 3.65 (s, 3H, N1–CH₃), 7.2–7.5 (m, 5H_{arom}), 8.2 (s, 1H, =CH) ppm; MS: m/z (%) = 394 (M⁺, 50), 366 (15), 289 (80), 261 (5), 233 (5), 204 (10), 176 (15), 135 (15), 106 (25), 96 (15), 77 (100), 67 (30), 51 (28), 42 (40).

7-Benzoylamino-1,3-dimethyl-9-(2,3,4,6-tetra-O-acetyl-β-Dglucopyranosyl)pyrimido[2,1-f]purine-1,2,3,4,6,9-

hexahydro-2,4,6-trione (**6**, C₃₁H₃₂N₆O₁₃)

To a solution of 3.7 g (0.01 mol) **3** in 0.01 mol aqueous potassium hydroxide in 6 cm³ distilled water, a solution of 4.15 g (0.011 mol) **5** was added. The mixture was stirred at room temperature until the reaction was judged complete by TLC [using CHCl₃/CH₃OH (90/10, ν/ν) as eluent, 15 h]. The mixture was evaporated under reduced pressure at 40°C and the residue was washed with distilled water to remove KBr. The product was filtered off, dried, and crystallized from dioxane.

Yield 5.6 g (80%); $R_{\rm f} = 0.42$; mp 280–282°C; IR: $\bar{\nu} = 3240$ (NH), 1750, 1710, 1666, 1590 (4CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 1.90-2.02$ (4s, 12H, 4COCH₃), 3.35 (s, 3H, N3–CH₃), 3.65 (s, 3H, N1–CH₃), 4.05 (m, 2H, 6', 6''–CH₂), 4.36 (m, 1H, 5'-H), 5.10 (t, 1H, 4'-H), 5.30 (t, J = 9 Hz, 1H, 2'-H), 5.70 (t, 1H, 3'-H), 6.20 (d, 1H, $J_{1',2'} = 10.5$ Hz, 1'-H), 7.2–7.5 (m, 5H_{arom}), 8.4 (s, 1H, =CH), 9.2 (s, 1H, NH) ppm.

1,3-Dimethylpyrimido[*2,1-f*]*purine-1,2,3,4-tetrahydro-2,4dione derivatives* (general procedure)

Compound 1 (0.2 g, 0.001 mol) was dissolved in 50 cm^3 dry *DMF* by heating, 1.4 g (0.015 mol) anhydrous potassium carbonate were added, followed by addition of 0.001 mol 7, 9, 11a, 11b, 15, 17, 19, 21, or 23. After being stirred under reflux for 10–15 h (TLC using ethyl acetate as eluent), the reaction mixture was concentrated *in vacuo*, poured into ice water, and neutralized with dilute HCl. The solid product precipitated, which was collected by filtration and recrystallized from the appropriate solvent.

8-Amino-1,3-dimethyl-6-(methylthio)-2,4-dioxo-1,2,3,4,8,9hexahydropyrimido[2,1-f]purine-7-carbonitrile

 $(\mathbf{8}, \mathbf{C}_{12}\mathbf{H}_{11}\mathbf{N}_7\mathbf{O}_2\mathbf{S})$

Reflux 15 h; yield 0.22 g (70%); $R_f = 0.19$; mp > 300°C (*DMF*); IR: $\bar{\nu} = 3364$, 3274 (2NH), 2225 (CN), 1697, 1639 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 2.6$ (s, 3H, SCH₃), 3.25 (s, 3H, N3–CH₃), 3.45 (s, 3H, N1–CH₃), 8.0 (s, 1H, NH), 9.6 (s, 1H, NH) ppm; MS: m/z (%) = 317 (M⁺, 100), 300 (10), 284 (10), 231 (20), 205 (5), 180 (3), 165 (30), 109 (25), 94 (10), 82 (35), 67 (35), 42 (35).

1,3-Dimethyl-6-(methylthio)-2,4,8-trioxo-1,2,3,4,8,9-hexahy-dropyrimido[*2,1-f*]*purine-7-carbonitrile* (**10**, C₁₂H₁₀N₆O₃S) Reflux 15 h; yield 0.21 g (65%); $R_{\rm f}$ =0.23; mp > 300°C (*DMF*); IR: $\bar{\nu}$ = 3448 (OH), 2217 (CN), 1705, 1655 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ = 2.7 (s, 3H, SCH₃) 3.20 (s, 3H, N3–CH₃), 3.50 (s, 3H, N1–CH₃), 9.6 (s, 1H, NH) ppm; MS: m/z (%) = 318 (M⁺, 40), 279 (10), 261 (15), 192 (10), 135 (5), 105 (15), 77 (20), 44 (100).

1, 3-Dimethyl-6, 8-diphenyl pyrimido [2, 1-f] purine-

2,4(1H,3H)-dione (12a, C₂₂H₁₇N₅O₂)

Reflux 10 h; yield 0.25 g (65%); $R_{\rm f} = 0.22$; mp > 300°C (*Et*OH); IR: $\bar{\nu} = 1702$, 1663 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ = 3.25 (s, 3H, N3–CH₃), 3.57 (s, 3H, N1–CH₃), 6.99 (s, 1H, 7-CH=), 7.47–8.37 (m, 10H_{arom}) ppm; MS: *m/z* (%) = 384 (M⁺ + 1, 30), 383 (M⁺, 100), 325 (51), 248 (62), 297 (24), 221 (12), 216 (15), 142 (22), 105 (79), 77 (59), 56 (13).

1, 3-Dimethyl - 2, 4-dioxo - 6, 8-diphenyl - 1, 2, 3, 4-tetrahydro-

pyrimido[2,1-f]purine-7-carbonitrile (**12b**, C₂₃H₁₆N₆O₂) Reflux 10 h; yield 0.22 g (55%); $R_{\rm f}$ =0.25; mp>300°C (*Et*OH); IR: $\bar{\nu}$ =2192 (CN), 1699, 1652 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ =3.38 (s, 3H, N3–CH₃), 3.45 (s, 3H, N1–CH₃), 7.52–8.07 (m, 10H_{arom}) ppm; MS: m/z(%) = 408 (M⁺, 23), 301 (10), 300 (19), 299 (100), 82 (38), 68 (21), 55 (24).

8-Amino-1,3-dimethyl-7-(4-nitrophenyl)pyrimido[2,1-f]purine-2,4(1H,3H)-dione (**14**, C₁₆H₁₃N₇O₄)

Reflux 10 h; yield 0.26 g (70%); $R_f = 0.21$; mp > 300°C (Dioxane/*DMF*); IR: $\bar{\nu} = 3360$, 3341 (2 NH), 1703, 1640 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.48$ (s, 3H, N3–CH₃), 3.52 (s, 3H, N1–CH₃), 7.35–8.35 (m, 5H_{arom}), 8.39 (s, 1H, 6-CH=), 9.2, 11 (s, 2H, NH₂) ppm; MS: m/z (%) = 367 (M⁺, 29), 366 (17), 306 (22), 234 (15), 194 (13), 179 (15), 152 (18), 127 (22), 104 (17), 99 (28), 82 (20), 59 (44).

1,3-Dimethyl-8-phenylpyrimido[2,1-f]purine-2,4(1H,3H)dione (16, $C_{16}H_{13}N_5O_2$)

Reflux 10 h; yield 0.18 g (60%); $R_f = 0.26$; mp > 300°C (*Et*OH); IR: $\bar{\nu} = 1703$, 1659 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.24$ (s, 3H, N3–CH₃), 3.44 (s, 3H, N1–CH₃), 7.54 (d, J = 7 Hz, 1H, 7-CH=), 7.57–7.87 (m, 5H_{arom}), 8.26 (d, J = 7 Hz, 1H, 6-CH=) ppm; MS: m/z (%) = 307 (M⁺, 66), 183 (33), 147 (33), 121 (53), 104 (100), 90 (53), 67 (35).

8-Amino-1,3-dimethyl-2,4-dioxo-6-phenyl-1,2,3,4-tetrahydropyrimido[2,1-f]purine-7-carbonitrile (**18a**, C₁₇H₁₃N₇O₂)

Reflux 12 h; yield 0.19 g (55%); $R_f = 0.19$; mp > 300°C (*Et*OH); IR: $\bar{\nu} = 3319$, 3260 (NH₂), 1704, 1637 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.38$ (s, 3H, N3–CH₃), 3.57 (s, 3H, N1–CH₃), 7.50–7.89 (m, 5H_{arom}), 8.36 (br, 2H, NH₂) ppm; MS: m/z (%) = 347 (M⁺, 25), 323 (32), 281 (33), 261 (29), 206 (62), 195 (49), 180 (18), 153 (27), 141 (8), 127 (32), 105 (11), 103 (23), 93 (30), 77 (78), 67 (67), 6 (31), 53 (49).

6-(4-Chlorophenyl)-1,3-dimethyl-2,4,8-trioxo-1,2,3,4,8,9hexahydropyrimido[2,1-f]purine-7-carbonitrile (**18b**, C₁₇H₁₂ClN₇O₂)

Reflux 12 h; yield 0.19 g (50%); $R_f = 0.18$; mp > 300°C; IR: $\bar{\nu} = 3320$, 3250 (NH₂), 1700, 1640 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.37$ (s, 3H, N3–CH₃), 3.42 (s, 3H, N1–CH₃), 7.50–7.88 (m, 4H_{arom}), 8.30 (br s, 2H, NH₂) ppm; MS: m/z (%) = 383 (M⁺+1, 29), 382 (M⁺, 41), 344 (41), 258 (79), 238 (73), 237 (85), 186 (41), 177 (38), 161 (100), 141 (52), 139 (47), 138 (61), 114 (38), 108 (35), 84 (67), 69 (61), 52 (44). 8-*Amino*-6-(2-*furyl*)-1,3-*dimethyl*-2,4-*dioxo*-1,2,3,4-*tetrahydropyrimido*[2,1-*f*]*purine*-7-*carbonitrile* (**18c**, C₁₅H₁₁N₇O₃) Reflux 12 h; yield 0.14 g (42%); R_f = 0.21; mp > 300°C (*DMF*); IR: $\bar{\nu}$ = 3396, 3210 (NH₂), 2201 (CN), 1698, 1649 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ = 3.47 (s, 3H, N3–CH₃), 3.51 (s, 3H, N1–CH₃), 7.10–7.70 (m, 3H, furan-H), 8.60 (br s, 2H, NH₂) ppm; MS: m/z (%) = 337 (M⁺,75), 256 (80), 249 (60), 210 (75), 199 (100), 140 (60), 128 (50), 121 (70), 110 (75), 108 (80), 86 (100), 83 (70), 68 (50), 56 (65).

8-Amino-1,3-dimethyl-2,4-dioxo-6-(2-thienyl)-1,2,3,4tetrahydropyrimido[2,1-f]purine-7-carbonitrile

 $(\textbf{18d},\,C_{15}H_{11}N_7O_2S)$

Reflux 12 h; yield 0.14 g (40%); $R_f = 0.20$; mp > 300°C (*DMF*); IR: $\bar{\nu} = 3326$, 3205 (NH₂), 2210 (CN), 1700, 1646 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.37$ (s, 3H, N3–CH₃), 3.43 (s, 3H, N1–CH₃), 7.00–7.40 (m, 3H, thiophene-H), 8.00 (br s, 2H, NH₂) ppm; MS: m/z (%) = 354 (M⁺+1, 9), 353 (M⁺, 5), 325 (30), 215 (7), 121 (7), 95 (11), 94 (100), 83 (10), 73 (9), 66 (10).

1,3-Dimethyl-2,4,8-trioxo-6-phenyl-1,2,3,4,8,9-hexahydro-

pyrimido[2,1-f]purine-7-carbonitrile (**20**, C₁₇H₁₂N₆O₃) Reflux 12 h; yield 0.14 g (40%); $R_{\rm f}$ =0.25; mp > 300°C (*Et*OH); IR: $\bar{\nu}$ =3334 (NH), 2203 (CN), 1703, 1654 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): δ =3.34 (s, 3H, N3–CH₃), 3.45 (s, 3H, N1–CH₃), 7.16–8.00 (m, 5H_{arom}), 8.10 (s, 1H, NH) ppm; MS: m/z (%) = 348 (M⁺, 23), 275 (25), 195 (84), 176 (4), 138 (14), 105 (24), 91 (23), 82 (20), 77 (19), 68 (15), 52 (8).

6-Acetyl-1,3,7-trimethyl-1H-imidazo[2,1-f]purine-

2,4(3H,8H)-dione (**22**, C₁₂H₁₃ N₅O₃)

Reflux 10 h; yield 0.18 g (65%); $R_f = 0.29$; mp > 300°C (*DMF*); IR: $\bar{\nu} = 3419$ (NH), 1700, 1652 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d_6, 200 MHz): $\delta = 2.27$ (s, 3H, CH₃), 2.44 (s, 3H, COCH₃), 3.17 (s, 3H, N3–CH₃), 3.47 (s, 3H, N1–CH₃), 8.71 (s, 1H, NH) ppm; MS: m/z (%) = 275 (M⁺, 92), 255 (52), 201 (68), 190 (48), 167 (60), 150 (72), 112 (60), 109 (60), 100 (44), 90 (80), 80 (48), 67 (100), 60 (60).

1,3-Dimethyl-7-phenyl-1H-imidazo[2,1-f]purine-

2,4(3H,8H)-dione (24 C₁₅H₁₃N₅O₂)

Reflux 10 h; yield 0.18 g (60%); $R_f = 0.25$; mp > 300°C (*EtOH*); IR: $\bar{\nu} = 3395$ (NH), 1698, 1644 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 3.40$ (s, 3H, N3–CH₃), 3.46 (s, 3H, N1–CH3), 7.31–8.08 (m, 5H_{arom}), 8.15 (s, 1H, CH=), 12.03 (s, 1H, NH) ppm; MS: m/z (%) = 295 (M⁺,18), 295 (9), 238 (4), 209 (4), 105 (100), 77 (61), 67 (5), 51 (20).

(6*E*)-1,3-Dimethyl-7-phenyl-1*H*-imidazo[2,1-*f*]purine-2,4,6(3*H*)-trione 6-[(4-methylphenyl)hydrazone] (**26**, C₂₂H₁₉N₇O₂)

A solution of 2.95 g (0.01 mol) **24** in 50 cm^3 ethanol was stirred with 1.4 g (0.01 mol) sodium acetate trihydrate for 15 min. The mixture was chilled in an ice bath at 0°C. While the solution was cooling, the 4-methylbenzene diazoni-

um chloride was prepared by the diazotization of 1.1g (0.01 mol) p-toluidine in $6 \text{ cm}^3 6M$ hydrochloric acid with $10 \,\mathrm{cm}^3$ cold 1M sodium nitrite solution in the usual way keeping the temperature below 5°C. The diazonium chloride solution was added to the reaction solution dropwise under stirring. The reaction mixture was left for 3 h in a refrigerator. The precipitated solid was filtered off, washed with water and ethanol, and dried. The product was recrystallized from 1,4-dioxane to give 26 as pure pale yellow crystals (TLC using ethyl acetate as eluent). Yield 0.29 g (70%); $R_{\rm f} = 0.19$; mp>300°C; IR: $\bar{\nu} = 3451$ (NH), 1696, 1643 (2CO) cm⁻¹; ¹H NMR (*DMSO*-d₆, 200 MHz): $\delta = 2.60$ (s, 3H, CH₃), 3.43 (s, 3H, N3-CH₃), 3.50 (s, 3H, N1-CH₃), 7.30-8.15 (m, 9H_{arom}), 8.35 (br s, 1H, NH) ppm; MS: m/z (%) = 414 (M⁺, 80), 413 (100), 307 (11), 282 (35), 234 (3), 207 (3), 195 (3), 106 (26), 94 (10), 77 (25), 67 (45), 53 (9); UV (methanol): $\lambda_{max}(\varepsilon) =$ 267 (18200), 461 (9800) nm (mol⁻¹ cm⁻¹).

Antimicrobial assay

Cultures of four fungal species namely Aspergillus fumigatus (AF), Penicillium italicum (PI), Syncephalastrum racemosum (SR), and Candida albicans (CA) as well as four bacterial species namely Staphylococcus aureus (SA), Pseudomonas aeruginosa (PA), Bacillus subtilis (BS), and Escherichia coli (EC) were used to investigate the antimicrobial activity of the compounds 3, 4, 8, 10, 12a, 12b, 16, 18c, 18d, and 26. The antimicrobial activity was assayed biologically using the diffusion plate technique. The latter technique was carried out by pouring a spore suspension of the fungal species (1 cm³ of sterile water contains approximately 108 conidia) or spreading bacterial suspension over a solidified malt agar medium. The layer is allowed to set for 30 min. A solution of the test compounds (1.0 g/cm^3) in *DMF* was placed onto sterile 5 mm filter paper discs and allowed to dry, then the discs were placed on the centre of the malt agar plate and incubated at optimum incubation temperature $28 \pm 2^{\circ}$ C. The fungicide Terbinfin and the bactericide chloramphenicol were used as standards under the same conditions. Measurements were considered after 72h for fungi and 24h for bacteria. The results are summarized in Table 1.

References

- 1. Corsano S, Scapiochi R, Strappaghetii G (1994) Arch Pharm (Weinheim) 327:411
- Baziard-Mouysset G, Rached A, Younes S, Tournaire C, Stigliani JL, Payard M, Yavo JC, Advenier C (1995) Eur J Med Chem 30:253
- 3. Sandoz AG (1992) Israeli IL 83:659; (1992) Chem Abstr 117:7957k
- 4. Gajewczyk L, Zejc A (1992) Acta Pol Pharm 49:61
- 5. Parrick J, Mehta LK (1993) J Heterocyclic Chem 30:323
- Bovy PR, Collins JT, Chamberlain TS, Gheng BK (1992) PCT Int ppl WO 92 07 852; Chem Abstr 117:131218j
- Kuzmenko VV, Pozharskii AF, Chemyshev AJ, Nanavyan IM (1987) Khim Geterotsikl Soedin 11:1551; (1988) Heterocyclic Compounds 240:94

- 8. Hesek D, Tegza M, Rybar A, Povazanec F (1989) Synthesis:681
- 9. Dreier A, Haller R (1987) Arch Pharm (Weinheim) 320:1004
- Nanavyan IM, Kuzmenko VV, Pozharskii AF, Klyuev AN (1987) Khim Geterotsikl Soedin 10:1398; (1988) Heterocyclic Compounds 206:97
- Karczmarzyk Z, Karolak-Wojciechowska J, Pawlowski M (1991) Acta Cryst 47:1902
- Karolak-Wojciechowska J, Pawlowski M (1990) J Crystallograph Spectrosc Res 20:477
- 13. Pawlowski M (1987) Pharmazie 42:371
- Pawlowski M, Buschauer A, Schunack W (1989) Arch Pharm (Weinheim) 322:447
- 15. Jin RH, Nisshjkubo T (1992) Tetrahedron Lett 33:6307
- Blythin DJ, Kamiinski JJ, Domalski MS, Spitler J, Solomon DM, Conn DJ, Shing-Chun W, Lehman LV, Bober LA, Chiu PJS, Watnick AS, Siegel MI, Hibert JM, Mc Phail AT (1986) J Med Chem 29:1099
- Conn DJ, Kamiinski JJ, Solomon DM, Mc Phail AT (1988) J Org Chem 53:3265
- Solomon DM, Kamiinski JJ (1989) USPat 4,816,458; Chem Abstr 111:153833j
- 19. Pawlowski M, Katlabi J, Rys B, Szneler E (1997) Pharmazie 52:279
- Pawlowski M, Katlabi J, Drabrzynska A, Duszynska B, Charakchieva-Minol S, Deren-Wesolek A, Tatarczynska E, Chojnacka-Wojcik E, Mokrosz MJ, Bojarski AJ (1999) Eur J Med Chem 34:167
- 21. Gatta F, Del Giudice MR, Borioni A, Mustazza C, (1994) J Heterocycl Chem 31:81
- 22. Harsanyi VK, Szebeni R, Korbonits D (1975) J Prakt Chem 317:745
- 23. Jin RH, Nishikubo T (1992) Tetrahedron Lett 33:6307
- 24. Lefebvro A, Rips R, Martine A, Lespagnol C (1985) J Heterocycl Chem 22:105
- Priimenko BA, Samura BA, Skulskaya EA, Troshin DA, Garmash SN, Milonova NP (1984) Khim Farm Zh 18:1456; (1985) Chem Abstr 103:22539e
- 26. Nosachenko VI, Kochergin PM, Steblyuk PN (1976) Khim Geterotsikl Soedin:1132; (1977) Chem Abstr 86:5414y
- 27. Hesek D, Tegza M, Rybar A, Povazanec F (1989) Synthesis:681
- Pawlowski M, Drabczynska A, Gorczyca M, Malecand D, Modzelewski J (1995) Pharmazie 50:453
- 29. Ueda T, Oh R, Nagai SI, Sakakibara J (1998) J Heterocycl Chem 35:135
- Da Settimo A, Da Settimo F, Marini AM, Primofiore G, Salerno S, Viola G, Via L, Magna SM (1998) Eur J Med Chem 33:685
- Eckstein M, Loson W (1968) Diss Pharm Pharmacol 20: 35; Abstract 69:43891h
- 32. Hino K, Irie A, Uno H (1975) Chem Pharm Bull 23:1696
- 33. Dreier A, Haller R (1986) Arch Pharm 320:999
- Da-Sellimo A, Prinofiore G, Marini AM, Dasetlimo F, La Motta C, Salerno S (1999) J Heterocycl Chem 36:639
- 35. Ueda T, Adachi T, Nagai S, Sakakibara J, Murata M (1988) J Heterocycl Chem 25:791

- Gatta F, Del Giudice MR, Borioni A, Borea PA, Dionisotti S, Ongini E (1993) Eur J Med Chem 28:569
- 37. Le Brun A (1971) Fr-Demande 2; 157:726; (1973) Abstract 79:126529d
- a) Mosselhi MAN (1990) PhD Thesis, Konstanz University, Germany; b) Jones JW, Robins RK (1960) J Am Chem Soc 82:3773
- 39. Mosselhi MAN (2002) Monatsh Chemie 133:1297
- 40. Zahran AM, El-Sharief AMS, El-Gaby MSA, Ammar YA, El-Said UH (2001) IL Farmaco 56:277
- Stanovnik B, Svete J, Tisler M, Zorz L, Hvala A, Simonic I (1980) Heterocycles 27:903
- 42. Gompper R, Topfl W (1962) Chem Ber 95:2861

- 43. a) Brunskill JSA, De A, Ewing DF (1978) J Chem Soc Perkin Trans I:629; b) Freemann F (1980) Chem Rev 80:329; c) Tornetta B, Scapini G, Guerrera F, Bernardini A (1970) Boll Seduta Accad Gioenia Sci Nat Catania 10:353; (1973) Chem Abstr 78:620n
- 44. Tseng SS, Epstein JW, Brabander HJ, Francisco G (1987) J Heterocyclic Chem 24:837
- 45. Elgemeie GEH, Attia AME, Fathy NM (1997) Nucleosides Nucleotides 16:485
- 46. Shawali AS, Mosselhi MAN, Tawfik NM (2001) J Org Chem 66:4055
- 47. Kralj L, Hvala A, Svete J, Golic L, Stanovnik B (1997) J Heterocycl Chem 34:247