Original Article

Synthesis and Antiplatelet Aggregation Activity Evaluation of some 2-Aminopyrimidine and 2-Substituted-4,6-diaminopyrimidine Derivatives

Marjan Esfahanizadeh^{*a,d*}, Shohreh Mohebbi^{*b*}, Behnam Dasht Bozorg^{*a*}, Salimeh Amidi^{*a*}, Ali Gudarzi^{*a*}, Seyed Abdolmajid Ayatollahi^{*a,c*} and Farzad Kobarfard^{*a,c,d**}

^aDepartment of Medicinal Chemistry, School of Pharmacy, Shahid Beheshti University of Medical Sciences, Tehran, Iran. ^bDepartment of Medicinal Chemistry, School of Pharmacy, Zanjan University of Medical Sciences, Zanjan, Iran. ^ePhytochemistry Research Center, Shahid Beheshti University of Medical Sciences, Tehran, Iran. ^dCentral Research Laboratories, Shahid Beheshti University of Medical Sciences, Tehran, Iran.

Abstract

A series of novel 2-aminopyrimidine and 2-Substituted-4,6-diaminopyrimidine derivatives have been synthesized and their antiplatelet aggregation activities were assessed against ADP and arachidonic acid-induced platelet aggregation in human plasma using light transmission aggregometry. Among the tested derivatives, compounds Ia, I_b, I_B and II₁₆ exhibited the highest antiplatelet aggregation activity (36.75, 72.4, 62.5 and 80 μ M). None of the compounds showed satisfactory activity against the aggregation induced by ADP but acceptable activities were observed against the aggregation induced by arachidonic acid. 2- aminopyrimidines were more active than 4,6- diaminopyrimidines in this respect.

Keywords: 2-aminopyrimidines; 2-Substituted-4,6-diaminopyrimidines; Antiplatelet aggregation.

Introduction

Platelets play an important role in maintaining cardiovascular integrity and in regulating the bleeding process by blood-clot formation (1). However, uncontrolled platelet aggregation is dangerous in arterial blockage and may lead to life threatening disorders (2). Antiplatelet agents are therefore considered as a significant tool in the treatment and/or prevention of cardiovascular thrombotic disease (3-5). Antiplatelet agents such as aspirin (acetylsalicylic acid), clopidogrel or ticlopidine and anticoagulants such as warfarin are currently two predominant groups of orally consumable drugs in standard therapeutic protocols for prophylaxis and treatment of venous thrombosis and reducing the risk of recurrent myocardial infarction (6-8).

Currently aspirin, which irreversibly inhibits cyclooxygenase I-mediated transformation of arachidonic acid to thromboxane A_2 (TXA₂), and the P_2Y_{12} antagonists clopidogrel and prasugrel, which selectively and irreversibly bind to the P_2Y_{12} ADP receptor are routinely used as antiplatelet agents (9, 10).

However there are still some serious limitations to these agents which include weak inhibition of platelet function (*eg.* aspirin) (11), slow onset of action (*eg.* clopidogrel) (12), variable response to treatment among the patients (*eg.* clopidogrel and aspirin) (11, 12) and high incidence of bleeding events which occur in both aspirin and clopidogrel drug therapy (13, 14).

^{*} Corresponding author:

E-mail: farzadkf@yahoo.com

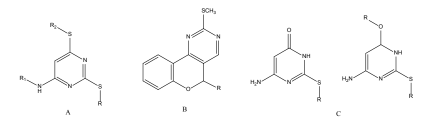


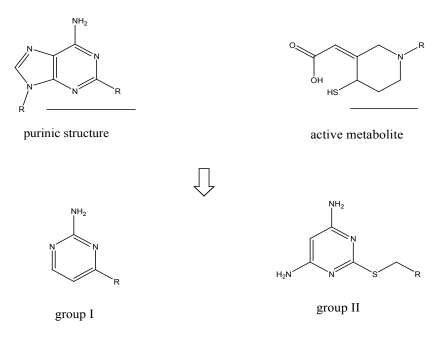
Figure 1. Active antiplatelet pyrimidine derivatives. A. 6-alkylamino-2,4-dialkyl(aryl)thiopyrimidines [7], R = Me, *n*-Pr, R1 = n-Pr, *i*-Pr, Ph, Bn, R2 = Bn, CH_2CH_2Ph ; B.Tricyclic pyrimidines [8], R = some amino substituents; c. Substituted 6-amino-2-mercaptopyrimidines [9], $R = CH_3$, CH_2CH_2NHBOC , CH_2CH_2OH , CH_2CF_3 , 2-Chlorobenzyl, *etc.* R1 = substituted arylsulfonyl moieties.

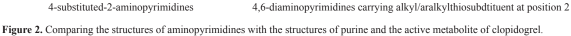
Considering the current situation, development of novel antiplatelet agents which are safe and effective is an urgent need (15).

Aminopyrimidine derivatives are an interesting group of compounds with various reported biological properties. Pyrimidine ring can be found in the structures of many important drugs such as nucleoside antibiotics, antibacterials and cardiovascular agents (16, 17).

Based on the hypothesis suggested by Cattaneo *et al.* (18), amino pyrimidine ring could be considered as a simplified form of the active metabolites of the thienopyridines and ATP derivatives. The active metabolites of thienopyridines have a simple monocyclic structure which implies that the presence of a bicyclic structure like that of a purine ring is not an absolute requirement for the affinity for the ADP receptor or so on platelet membrane.

A group of amino pyrimidine derivative with thioether substituents has been synthesized and evaluated by Cattaneo *et al.* The compound showed satisfactory anti platelet aggregation effects when ADP had been used for aggregation induction (18). Based on the mentioned reports and in order to investigate the capability of amino pyrimidine derivatives in inhibition of platelet aggregation pathways, we synthesized two groups of amino pyrimidines with the following structures:





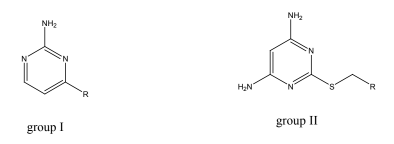


Figure 3. Chemical structure of the synthesized compounds.

Comparing the activity of these compounds in inhibition of platelet aggregation induced by ADP and arachidonic acid will provide some insights into the structure activity relationship of these compounds.

Chemistry

The synthetic procedures for groups I and II are illustrated in Figures 4 and 5.

Group I: Methyl ketones (1a-h) were allowed to react with dimethylformamidedimethylacetate (DMF-DMA) to produce 3-(dimethylamino)-1-aryl-2en-1-ones (2a-h). These intermediates could be then condensed with guanidine HCl to obtain the corresponding amino pyrimidine ring systems (19).

Group II: As it is shown in Figure 5 the intermediate II_3 (4,6-diaminopyrimidine-2-thiol) was obtained by the reaction of thiourea and malononitrile in absolute ethanol as the solvent. Subsequent reaction of compound II_3 with various alkylhalides at room temperature afforded compounds II_{4-25} in good yields.

Structure confirmation of the synthesized intermediates and the final products was performed using IR, NMR and Mass spectrometry (20).

Experimental

General

All evaporations were carried out in vacuo with a rotary evaporator. Melting points (°C) were determined by capillary method on an electrothermal melting point apparatus. Infrared spectra were recorded as thin films of KBr plates with U_{max} in inverse centimeters. Nuclear magnetic resonance spectra for proton (1H NMR) were recorded on a Bruker DRX-Avance (500 MHz) spectrometer. Chemical shift values are expressed in ppm (parts per million) relative to tetramethylsilane (TMS) as internal standard; s: singlet, d: doublet, dd: double doublet, t: triplet, q: quartet, m: multiplet, br s: broad singlet. Thin layer chromatography (TLC) was performed on Whatman SilG/UV₂₅₄ silica gel plates with fluorescent indicator and the

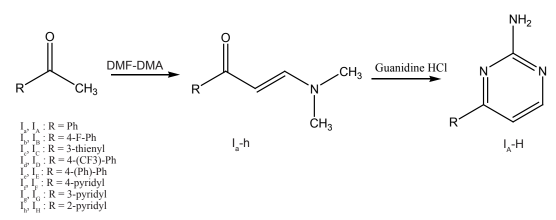


Figure 4. Compounds (I_{a-h}) and (I_{A-H}) synthesis scheme. Reagent and conditions: (a) DMF, reflux, 24 h; (b) NaOCH₃, Isopropanol, reflux, 48 h.

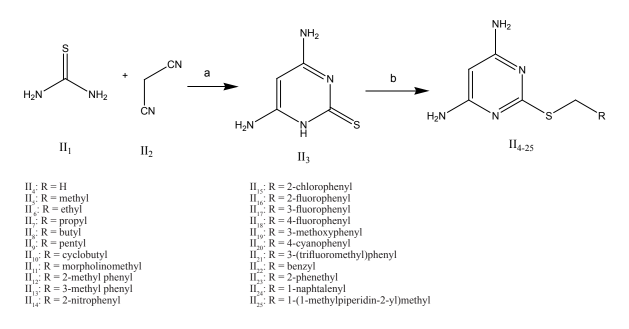


Figure 5. Compounds (II₄₋₂₅) synthesis scheme. Reagents and conditions: (a) EtONa, reflux, 3 h; (b) NaOH 0.1 M, CH₃OH, R-CH₂-X (X = Cl, Br), rt, 18 h10).

spots were visualized under 254 and 366 nm illumination. Mass analyses were performed on an Agilent 6400 series equipped with an electrospray (ESI) ionization interface (drying gas adjusted at 300 °C, nebulizing gas flow at 12 L/min). All the compounds were analyzed for C, H, N and S on a Costech model 4010 and agreed with the proposed structures within $\pm 0.4\%$ of the theoretical values.

General procedure for the preparation of compounds $(I_{a,b})$

The detailed description of the methods used for preparation of compounds I to I and compounds I_A to I_H is reported in reference 19. Briefly to a solution of acetophenone (84 mmol) in DMF (16 mL), was added DMF-DMA (84 mmol) and the solution was heated under reflux for 24 h. Brine (25 mL) was added to the reaction mixture after cooling and the reaction mixture was then extracted with CH_2Cl_2 (3 × 25 mL). The combined organic fractions were dried over anhydrous Na₂SO₄ and the solvent was evaporated under reduced pressure. The residue was dissolved in EtOAc (12 mL), followed by addition of n-hexane (100 mL). The precipitate thus obtained was filtered and dried to give the pure product I as

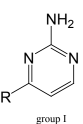
a yellow solid (19).

General procedure for the preparation of compounds (I_{A-H})

To a solution of compounds I_a (2.68 mmol) in isopropanol (13.5 mL) were added sodium methoxide (10.7 mmol) and guanidine hydrochloride (4.02 mmol) and the solution was heated under reflux for 48h. Distilled water (25 mL) was added to the reaction mixture after cooling and the mixture was then extracted with EtOAc (3 × 15 mL). The combined organic fractions were dried over anhydrous Na₂SO₄ and the solvent was evaporated under reduced pressure. The residue was dissolved in EtOAc (2 mL), followed by addition of *n*-hexane (25 mL). The precipitate thus obtained was filtered and dried to give I_{A-H} .

Representative procedure for the preparation of compounds (II_{4-25})

The detailed description of the methods used for preparation of compounds II_4 to II_{25} is reported in reference 20. Briefly, to a solution of 4,6-diaminopyrimidine-2-thiol (II₃, 1.4 mmol) in methanol, was added alkyl halide (3.5 mmol) under basic conditions (NaOH 0.1 M, 14 mL). The mixture was then stirred for 18h at Table 1. Antiplatelet activity of group I derivatives.



Compound	R	A.A IC ₅₀ MM	ADP %inhibition
Ia (1.25 mM)	ph	36.75	45.2
I _A (0.75 mM)	ph	544	29
Ib (1 mM)	4-F- ph	72.4	28.6
I _B (2.5 mM)	4-F- ph	62.5	53
I _c (1 mM)	3-thienyl	1000	35.5
I _c (1.6 mM)	3-thienyl	>1000	57.3
I _d (1.3 mM)	4-(CF ₃)- ph	>1000	24.3
I _D (0.5 mM)	4-(CF ₃)- ph	340	42.5
Ie (1.3 mM)	4-(ph)- ph	>1000	24.3
I _E (1 mM)	4-(ph)- ph	>1000	50
If (1 mM)	4-pyridyl	>1000	1
$I_F(1 \text{ mM})$	4-pyridyl	ND	ND
Ig (2.5 mM)	3-pyridyl	1000	35
I _G (0.5 mM)	3-pyridyl	192	39.5
I _h (1.25 mM)	2-pyridyl	752	27.1
I _H (1.25 mM)	2-pyridyl	>1000	44
Indomethacin		3.0	
Aspirin		30.3	

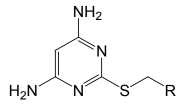
room temperature. After removing the solvent under reduced pressure, the residue was washed by water and the precipitate was collected as a solid. All compounds were obtained in acceptable purity and no further purification was needed (20).

Results and Discussion

All the synthesized compounds were screened for their effects on human platelet aggregation induced by arachidonic acid and ADP using light transmission aggregometry. IC_{50} was determined as the concentration of the test compounds that exhibit platelet aggregation

by 50%. The *in-vitro* antiplatelet activity of the synthesized derivatives is listed in Tables 1 and 2.

Comparing the activities of the two aminopyrimidine groups indicates that none of the compounds showed satisfactory activity against the aggregation induced by ADP. Therefore it could be concluded that the compounds do not interfere with ADP receptors on platelet membrane. However, acceptable activities were observed in both groups against the aggregation induced by arachidonic acid. This is not in agreement with the report by Cattaneo *et al.* who introduced a group of aminopyrimidines as active platelet aggregation Table 2. Antiplatelet activity of group II derivatives.



group II

Compound (In 1 mM)	R	A.A IC ₅₀ MM	ADP % inhibition
II ₄	Н	>1000	10
II ₅	Methyl	>1000	33.8
II_{6}	Ethyl	500	42
II ₇	Propyl	ND	ND
II ₈	Butyl	500	42
II ₉	Pentyl	>1000	53
II ₁₀	Cyclobutyl	>1000	49
II ₁₁	Morpholinomethyl	>1000	ND
II ₁₂	2-methylphenyl	213	38.3
II ₁₃	3-methylphenyl	220	10
II_{14}	2-nitrophenyl	209	20
II ₁₅	2-chlorophenyl	>1000	14.7
II ₁₆	2-fluorophenyl	80	23
II ₁₇	3-fluorophenyl	700	49
II ₁₈	4-fluorophenyl	214	24.3
II ₁₉	3-methoxyphenyl	>1000	8
II ₂₀	4-cyanophenyl	>1000	28
II ₂₁	3-(trifluoromethyl)phenyl	>1000	50
II ₂₂	Benzyl	ND	ND
II ₂₃	2-phenethyl	760	70
II_{24}	1-naphthalenyl	1000	51.9
II ₂₅	1-(1-methylpiperidin-2-yl)methyl	>1000	95
Indomethacin		3.0	
aspirin		30.3	

inhibitors which interfere with ADP receptors.

Comparing the overall results obtained for aminopyrimidines I and II indicates that 2-aminopyrimidines (I) were more active than 4,6-diaminopyrimidines (II). Only compound 16 in group II showed satisfactory IC_{50} (80 µM).

Among the 2-aminopyrimidines group (I), on the other hand, a few compounds (I_a , I_b , I_B and I_G) showed good activities (36.75, 72.4,

62.5 and 192 μ M). Interestingly, compounds with fluorine substituent on phenyl ring (I_b , I_B) were among the most active compounds.

Global physicochemical properties for compounds I_{a-h} , I_{A-H} and II_{4-25} were calculated using Chemdraw Ultra, Chem3D Ultra version 8.0 and Hyper Chem professional and the results are presented in Tables 3 and 4.

Efforts to find a relationship between these

Compound	ClogPA	Р ^в	Vc	SA ^D	D E
Ia	2.2974	20.85	606.6	388.66	2.669
I _A	1.774	20.03	549.5	259.2	1.319
Ib	2.516	24.43	632.5	398.5	3.181
I _B	1.926	22.06	553.45	357.2	1.122
I _c	2.0739	20.38	581.23	379.5	2.56
I _c	1.4697	19.55	521.3	342.54	0.778
I _d	3.3127	22.42	683.67	430.8	3.844
I _D	2.6730	21.59	620.86	395.87	2.208
Ie	4.185	30.5	811.04	495.06	2.708
I_E	3.66	29.69	747.46	458.7	1.548
If	1.1964	20.14	592.2	382.9	3.2
I_F	0.409	19.32	529.56	344.89	1.194
Ig	1.1964	20.14	591.97	383.02	1.283
I_{G}	0.1997	19.32	529.45	345.85	1.944
I _h	1.5964	22.95	611.69	389.11	1.914
I _H	0.409	19.32	535.7	351.3	1.3

Table 3. Global	physicochemical	properties for	compounds group I.

^AClogP were calculated by using Chem Draw Ultra version 8.0. ^BPolarizability values were calculated by using Hyper Chem Professional. ^D Surface area values were calculated by using Hyper Chem Professional. ^D Durface area values were calculated by using Hyper Chem Professional. ^CDipole (debye) values were calculated by using Chem3D Ultra version 8.0.

Compound	ClogPA	Рв	Vc	SAD	\mathbf{D}^{E}
II ₄	0.866	16.55	487.68	331.52	3.1
II_5	1.395	18.39	542.6	362.6	2.926
II_6	1.924	20.22	596.38	394.4	2.933
II ₇	2.453	22.06	650.7	423.37	2.895
II_8	2.982	23.89	703.83	455.17	2.9047
II_9	3.511	25.73	749.73	477.11	2.884
II_{10}	2.398	23.12	653.14	414.09	2.898
II ₁₁	0.8381	26.94	745.2	461.24	3.168
II_{12}	2.883	28.05	744.66	467.25	3.084
II ₁₃	2.933	28.05	754.32	472.14	2.902
II_{14}	2.097	32.61	767.72	477.94	4.137
II ₁₅	3.147	28.14	729.07	460.65	41.994
II ₁₆	2.577	26.12	706.53	448.67	1.818
II ₁₇	2.577	26.12	711.03	452.9	3.835
II_{18}	2.577	26.12	711.7	453.5	4.995
II ₁₉	2.353	28.68	771.85	485.4	1.774
II_{20}	1.867	28.06	760.45	479.2	6.326
II ₂₁	3.317	27.77	779.18	489.7	4.5
II ₂₂	2.963	28.05	754.7	477.364	3.067
II ₂₃	3.342	29.88	809.23	507.6	3.145
II ₂₄	3.608	33.48	826.6	509.25	2.885
II ₂₅	2.397	29.47	799.19	477.84	1.924

Table 4.Global	physicochemical	properties for com	pounds group II.

^AClogP were calculated by using Chem Draw Ultra version 8.0. ^BPolarizability values were calculated by using Hyper Chem Professional. ^CMolecular volume values were calculated by using Hyper Chem Professional. ^DSurface area values were calculated by using Hyper Chem Professional. ^CDipole (debye) values were calculated by using Chem3D Ultra version 8.0.

physicochemical parameters and anti platelet aggregation activity of the compounds did not result in a clear correlation. This could be due to the fact that the compounds are different in their access to their targets in the platelet aggregation pathway induced by arachidonic acid.

Further mechanistic studies are needed to clarify the mechanism of antiplatelet activity of these compounds.

References

- Zia-Ul-Haq M, Ahmad Shahid S, Ahmed S, Ahmad Sh, Qayum M and Khan I. Antiplatelet activity of methanolic extract of Grewiaasiatica L. leaves and Terminallachebula Retz Fruits. *J. Med. Plants Res.* (2012) 6: 2029-2032.
- (2) Mathers C, Fat DM and Boerma JT. World Health Organization. *The Global Burden of Diseases: 2004 Update*. Geneva (2008).
- (3) White HD. Oral antiplatelet therapy for a therothrombotic disease: Current evidence and newbdirections. Am. Heart J. (2011) 161: 450-461.
- (4) Latib A, Ielasi A, Ferri L, Chieffo A, Godino C, CarlinoM, Montorfano M and Colombo A. Aspirin intoleranceand the need for dual antiplatelet therapy after stentimplantation: A proposed alternative regimen. *Int. J. Cardiol.* (2013) 165: 444-447.
- (5) Reiter RA and Jilma B. Platelets and new antiplateletdrugs. *Therapy* (2005) 2: 465-502.
- (6) Ilas J, Jakopin Z, Borstnar T, Stegnar M and KikeljD. 3,4-Dihydro-2H-1,4-benzoxazine derivativescombining thrombin inhibitory and glycoprotein IIb/IIIa receptor antagonistic activity as a novel class ofantithrombotic compounds with dual function. J. Med. Chem. (2008) 51: 5617-5629.
- (7) De Meyer SF, Vanhoorelbeke K, Broos K, Salles II and Deckmyn H. Antiplatelet drugs. *Br. J. Haematol.* (2008) 142: 515-528.
- (8) AmidiS, Kobarfard F, Moghaddam AB, Tabib K and SoleymaniZ. Electrochemical synthesis of novel 1,3-indandione derivatives and evaluation of their antiplatelet aggregation activities. *Iran. J. Pharm. Res.* (2013) 12: 91-103.
- (9) Maree AO and Fitzgerald DJ. Variable platelet response to aspirin and clopidogrel in atherothrombotic disease.

Circulation (2007) 115: 2196-2207.

- (10) Meadows TA and Bhatt DL. Clinical aspects of platelet inhibitors and thrombus formation. *Circ. Res.* (2007) 100: 1261-1275.
- (11) Patrono C, Coller B, Fitz Gerald GA, Hirsh J and Roth G. Platelet-active drugs: the relationships among dose, effectiveness, and side effects. *Chest.* (2004) 126: 234-264.
- (12) Bassand JP. Unmet needs in antiplatelet therapy. *Eur. Heart J.* (2008) 10: 3-11.
- (13) Guthrie R. Review and management of side effects associated with antiplatelet therapy for prevention of recurrent cerebrovascular events. *Adv. Ther.* (2011) 28: 473-482.
- (14) Chan YC, Valenti D, Mansfield AO and Stansby G. Warfarin induced skin necrosis. *Br. J. Surg.* (2000) 87: 266-272.
- (15) Liu G, Xu J, Chen N, Zhang S, Ding Z and Du H. Synthesis of N6-alkyl(aryl)-2-alkyl(aryl) thioadenosines as antiplatelet agents. *Eur. J. Med. Chem.* (2012) 53: 114-123.
- (16) Tozkoparan B, Ertan M, Kelicen P and Demirdamar R. Synthesis and anti-inflammatory activities of some thiazolo[3,2-a]pyrimidine derivatives. *Farmaco*. (1999) 54: 588-593.
- (17) Clark J, Shahhet MJ,Korakas D and Varvounis
 G. Synthesis of thieno[2,3-d]pyrimidines from 4,6-dichloropyrimidine-5-carbaldehydes. J. Heterocyclic Chem. (1993) 30: 1065-1072.
- (18) Crepaldia P, Cacciari B, Bonache MC, Spalluto G, Varani K, Borea PA, Kugelen IV, Hoffman K, Pugliano M, Razzari C and Cattaneo M. 6-amino-2-mercapto-3H-pyrimine-4-one derivatives as new candidates for the antagonism at the P2Y₁₂ receptors. *Bioorg. Med. Chem. J.* (2009) 17: 4612-4621.
- (19) Mohebbi Sh, Shirazi FH, Sharifnia SH and Kobarfard F. Introducing synthesis route-based hit identification approachas a tool in medicinal chemistry and its application in investigating the antiproliferative and antimicrobial effects of 2-aminopyrimidine derivatives. *Int. J. Drug Discov.* (2011) 3: 78-87.
- (20) Mohebbi Sh, Falcón-Pérez JM, González E, Millet O, Mato JM and Kobarfard F. Synthesis, dihydrofolate reductase inhibition, anti-prolifrative testing and saturation transfer difference ¹H-NMR study of some new 2-substituted-4,6-diaminopyrimidine derivatives. *Chem. Pharm. Bull.* (2012) 60: 70-78.

This article is available online at http://www.ijpr.ir