

Benzothiazole: As an Antiviral Agent

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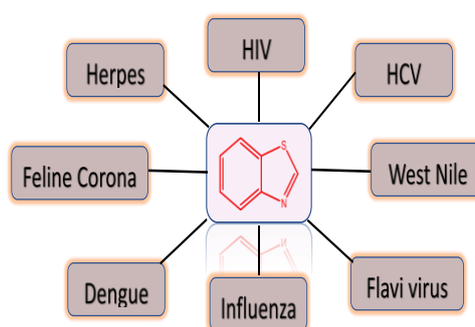
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Abstract: The virus is a microorganism that uses the machinery of the host to multiply. At present, there are various species of viruses known to us that are dangerous for the health of human beings. One of such viruses has destroyed many lives nowadays and that is a coronavirus. Such other viruses like Human Immunodeficiency Virus, Poliovirus, etc. destroy one's capability to survive normally. As science progresses, we invent many antiviral drugs as per the type of virus. There are many antiviral drugs available to treat viral infections. From them, benzothiazole derivatives are potent antiviral agents. Researchers continuously work on benzothiazole moiety to get more effective benzothiazole derivatives that can be used as antiviral agents. This review article gives information about various benzothiazole derivatives that act against the various viruses as antiviral agents, the structure-activity relationship of benzothiazole as an antiviral agent, various schemes to synthesize benzothiazole derivatives as an antiviral agent as well as includes various methods to evaluate the antiviral activity of novel synthetic compounds against specific viruses.

Keywords: Benzothiazole antiviral derivatives; schemes for synthesis; structure-activity relationship; in vitro methods to evaluate antiviral activity

Graphical Abstract



1. Introduction

The Virus is very tiny infectious agent that replicate only inside the living cells of an organism. Viruses infect all types of plants, animals and microorganisms also, like bacteria etc. [1] Viral infections are considered to be one of the major threats to the health of human being. Virus infections take place due to globalization and unexpected climate change.[2] We are informed about just about 260 varieties of viruses, but the unknown varieties of viruses are responsible for 99.9% of total infection cases. These viruses come to the picture

when they show some symptoms to the host.[3] Despite of the development of many molecules as antiviral, they are unable to satisfy the requirement criteria to treat the viral infection and drug resistance of current viruses. That's why, it's still need of newer vaccines, diagnostic agents and antiviral molecules.[4] Because, benzothiazole is such a versatile moiety, it shows many biological activities including antiviral effect against various species of viruses[5]. Because of feasible physical and chemical properties of benzothiazole moiety, many researchers have tried to synthesize various benzothiazole derivatives that shows potent antiviral effects against various strain of viruses[6].

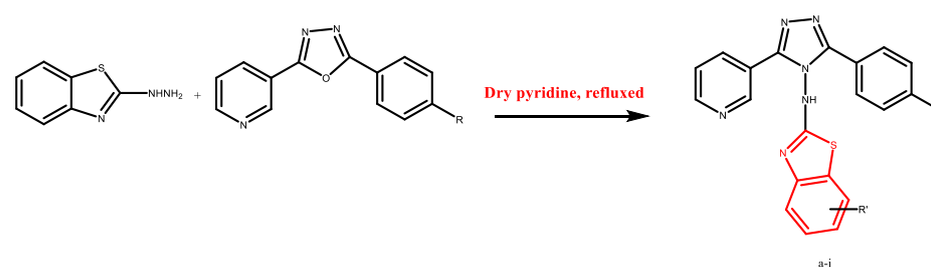
Structurally benzothiazole is a fusion of two aryl rings; benzene and thiazole. As thiazole bears nitrogen and sulfur moiety, benzothiazole derivatives successfully binds to viruses and gives antiviral activity. Many articles are available for reference to develop newer antiviral agents, but this review article includes various novel synthesized antiviral compound bearing benzothiazole moiety, pathway to synthesize benzothiazole based antiviral agents, structure-activity relationship of benzothiazole as antiviral agents and in vitro and in vivo methods to evaluate antiviral activity of novel synthetic compounds. That's why it is a unique article containing all the information to guide a researcher for synthesis of benzothiazole based antiviral agents.

2. Pathway to Synthesize Antiviral Drugs Bearing Benzothiazole Moiety

Navin B. Patel and his colleagues designed the above pathway to synthesize benzothiazole derivatives having antiviral activity. These compounds were active against Human Immuno Deficiency Virus. 2-aminobenzothiazole in dry pyridine was refluxed with 2-(4-nitrophenyl)-5-(pyridin-3-yl)-1,3,4-oxadiazole or 2-(4-chlorophenyl)-5-(pyridin-3-yl)-1,3,4-oxadiazole to get the respective benzothiazole derivatives.[7] (Scheme 1)

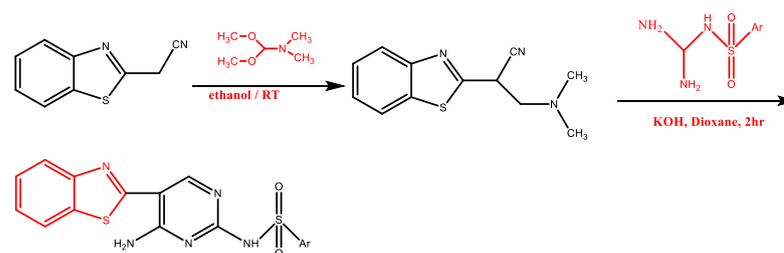
Rasha et al. synthesized benzothiazole derivatives by scheme 3. Benzothiazole-2-yl-acetonitrile was allowed to react with N,N-dimethylformamide dimethyl acetal in ethyl alcohol at room temperature for 10 min to get the 2-(benzo[d]thiazol-2-yl)-3-(dimethylamino)acrylonitrile. This intermediate was further reacted with N-arylsulfonated guanidine in presence of potassium hydroxide and dioxane for 2 hours. The resulted compounds were (4-amino-5-(benzo[d]thiazol-2-yl)pyrimidin-2-yl)-arylsulfonamides that show potent antiviral activity against Herpes Simplex Virus. They are also Hsp90 α inhibitors with broad spectrum antiviral activity. [8] (Scheme 2)

Scheme 1



R= NO₂, Cl; R' = a) 6-F, b) 6-Br, c) 6-NO₂, d) 6-CH₃, e) 6-OCH₃, f) 6-Cl, g) 4-CH₃, h) 4-NO₂ i) 5-Cl, 6-Cl, j) 4-Cl

Scheme 2



a) Ar = C₆H₅ b) Ar = 4-CH₃-C₆H₄

incorporation of [3H]-UTP by recombinant HCV RNA polymerase NS5BDC21 (genotype 1b) on a homopolymeric RNA primer/template. The non-nucleoside inhibitor aurintricarboxylic acid (ATA) was used as reference compound in the enzymatic assay. This compound has been shown to inhibit NS5B both in vitro and in replicon assays, through binding to the benzothiadizine allosteric pocket.[10]

Girijavallabhan et al. have designed some HCV replication inhibitors bearing benzothiazole moiety. Compound (3) was screened for its HCV replication inhibition ability and it was found the potent compound as an antiviral compound.[11]

3.2. Anti-Herpes Virus Agents

Human cytomegalo virus is under the category of beta herpes virus. Once the person is infected with this particular virus, the virus remains for whole life into that person's body. It does not affect to the healthy human but it shows symptoms in pregnant lady or the persons with weak immunity.[12] Then the treatment becomes necessary.

The substituted benzothiazole derivatives (4) and (5) were prepared by reacting lactam with respective substituted bromobenzothiazole under Cu catalysis using modified Goldberg conditions by Alan et al. compounds (4) and (5) were tested against HCMV virus using modified ELISA technique. The compounds show potent antiviral activity against HCMV - Human Cytomegalovirus.[13]

Hatem et al. have synthesized (1Z,2E)-2-(2-(benzo[d]thiazole-2-carbonyl)hydrazineylidene)-N'-(4-chlorophenyl)-N-hydroxypropanehydrazonamide and other benzothiazole derivatives having antiviral activity. Benzoyl hydrazine was reacted with 2-oxo-N-arylpropanehydrazonoyl chlorides by refluxing with ethanol. The reaction was resulted in corresponding hydrazonoyl chloride which was further reacted with benzene sulphinate to have corresponding sulphones. Reaction of some hydrazonoyl chloride with hydroxylamine hydrochloride in presence of potassium carbonate, resulted in N-hydroxy-2-(2-(benzothiazole-2-carbonyl)hydrazono)-N9-(4-aryl)propanehydrazonamide (6). This compound is active against Herpes simplex type 1 virus. (HSV-1)[14]

3.3. Anti-Dengue Virus Agents

The main cause of dengue is flaviviridae virus with the carrier mosquito. This is a single stranded RNA virus who infect almost 50 million people every year. Now a days no specific agents are available to treat dengue but the effective molecules directly target the viral structural proteins.[15] Various benzothiazole derivatives have been screened for their anti-dengue activity and some of them shows potent activity against dengue.

Compound (7) was designed by Halim et al. for screening against dengue virus. Novel DENV NS-3 helicase inhibitor from zinc database was confirmed having antiviral activity against dengue virus by computational modelling techniques as well as in vitro and in vivo biological assays.[16]

3.4. Anti-HIV Virus Agents

Human Immunodeficiency Virus is the most dangerous virus in the virus family. The resultant disorder is Auto Immunodeficiency Disorder. For the treatment of AIDS, no any specific treatment is available till now. The current therapy against AIDS, is based on six of categories drugs: nucleoside/nucleotide reverse transcriptase inhibitors (NRTIs/NtRTIs); nonnucleoside reverse transcriptase inhibitors (NNRTIs); protease inhibitors (PIs); cell entry inhibitors [fusion inhibitors (FIs) and co-receptor inhibitors (CRIs)]; and integrase inhibitors (INIs).[17] HAART (Highly active anti-retroviral therapy) is beneficial but it has very high cost and severe toxicity. But some benzothiazoles are analyzed for their anti-HIV activity which may be at low cost and higher potency.

2-(4-(1-benzyl-2-ethyl-4-nitro-1H-imidazol-5-yl)piperazin-1-yl)benzo[d]thiazole (15) was synthesized and analyzed by Yaseen et al. for their anti-HIV activity. 1-benzyl-5-

bromo-2-ethyl-4-nitro-1H-imidazole was reacted with 1-(1-benzyl-2-ethyl-4-nitro-1H-imidazol-5-yl)piperazine it will result in the targeted compound. The resulted compound (8) was assumed to act as non-nucleoside reverse transcriptase inhibitor (NNRI).[18]

Al-Masoudi et al. have designed a series of benzothiazole derivatives. The compounds were tested for in vitro activity against HIV-1 and HIV-2 in human T-lymphocyte (MT-4) cells based on the MTT assay. From which (3-(Benzothiazol-2-ylmethyl)-6-methoxy-2-methyl-1H-indol-1-yl)(4-chlorophenyl)methanone (9) was found to be active against strain of human immunodeficiency virus.[19]

3.5. Anti-Influenza Virus Agents

Influenza virus is commonly known as “flu” and main cause of this infection is influenza A or influenza B virus. The machinery that the virus contains is single stranded RNA virus. It affects the upper respiratory tract therefore the symptoms involve common cold and fever. 36 benzothiazole derivatives that act against influenza virus are 1-(benzo[d]thiazol-2-yl)-3-methylurea³⁷ (10) and 9-((5-(benzo[d]thiazol-2-ylamino)pentyl)oxy)-10-methoxy-[1,3]dioxolo[4,5-g]isoquinolino[3,2-a]isoquinolin-7-ium (11) [20]

4. Structure Activity Relationship of Benzothiazole Derivatives as Antiviral Agents

- Amine or amido linkage at 2nd position of benzothiazole gives anticancer activity of compound.
- Second position of benzothiazole is active to attach substituents.
- Methyl group substitution at 5th or 6th position of benzothiazole, increase potency of antiviral compounds.
- Aryl moieties like pyrazole, pyridine, phenyl, imidazole, benzothiazole, thiazole etc at 2nd position gives antiviral activity of the compound. Directly attached or through amine or amide linkage, aryl moiety at 2nd position of benzothiazole gives potent antiviral compounds.
- 4th position of benzothiazole is also important for substitution in 2-aminobenzothiazoles to derive antiviral moieties.

5. Methods to Evaluate Antiviral Activity of Benzothiazole Derivatives

The antiviral activity of novel synthetic compound can be measured by the four following methods in vitro.

5.1. Inhibition of Virus Induced Cytopathic Effect: [21,22]

A quantal assay can be used to determine effectiveness of those viruses that induce cytopathic effect but do not cause plaque reduction. Here is the procedure to follow the assay method: Prepare a series of one fourth of cell cultures in culture tubes which are infected with a constant dose of 100 TCID₅₀ (Median Tissue Culture Infectious Dose). The series should be kept at 37°C for 1-2 hr. Add antiviral agent with maintenance media to that series after completion of 1-2 hr. Concentration range of antiviral agent should be from minimal dose that doesn't show any antiviral activity to the maximum dose of inhibition. Virus induced cytopathic effect will be recorded each day until all the samples and blank cultures show cytopathic effect. ED₅₀ that is the 50% of the effective dose of antiviral drug is the concentration that inhibits cytopathic effect in half of the culture tubes.

5.2. Plaque Reduction Assay: [23]

This method is applicable to all viruses that form plaque in suitable cells. This method is performed in corresponding cell monolayers infected with a constant concentration of virus depending on the size of monolayer. The monolayers are kept at 37°C for 1-2 hr. and after those nutrients and antiviral agent with 1-2% methylcellulose is added. The infected culture is kept for rest for incubation for a respective period of time for different species of viruses. At the end of incubation period, all the cultures are stained for examination of

plaque number. Plaque numbers in the culture without antiviral agent and with different concentration of antiviral agents are compared for result.

5.3. Virus Yield Reduction Assay: [24,25]

This method is used when long time determination of antiviral agent becomes necessary. Here, the cultures are infected with a fixed dose of virus and kept for 2 hrs. at 37 C then the unabsorbed viruses are removed by washing with Hanks' Balanced Salt Solution and the antiviral drug of various concentrations are added. After incubation period, the cultures are tested for total virus yield. ED90 (Drug concentration 90% of the virus yield) in comparison with virus control is determined from dose response curves.

5.4. Assay Systems Based on Measurement of Specialized Functions and Viral Products: [26-30]

Certain viruses do not produce plaque or cytopathic effect but perform some specific functions so we can use them as evaluation parameter. The specific functions could be hemagglutination, hemadsorption, extent of viral replication, reduction of virus specific polypeptides, synthesis of viral nucleic acids, reverse transcriptase activity, by dose response curve etc. from these all measurements, one can derive ED50 value of antiviral agent and can compare it with virus control to evaluate antiviral activity of the compound.

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Conflicts of Interest: No conflict of interest.

References

1. Tripathi K.D. Essentials of medical pharmacology. JP Medical Ltd; 2013 Sep 30.
2. Cavicchioli R, Ripple WJ, Timmis KN, Azam F, Bakken LR, Baylis M, Behrenfeld MJ, Boetius A, Boyd PW, Classen AT, Crowther TW. Scientists' warning to humanity: microorganisms and climate change. *Nature Reviews Microbiology*. 2019 Sep;17(9):569-86.
3. Carroll D, Watson B, Togami E, Daszak P, Mazet JA, Chrisman CJ, Rubin EM, Wolfe N, Morel CM, Gao GF, Burci GL. Building a global atlas of zoonotic viruses. *Bulletin of the World Health Organization*. 2018 Apr 1;96(4):292.
4. Hassan MZ, Osman H, Ali MA, Ahsan MJ. Therapeutic potential of coumarins as antiviral agents. *European journal of medicinal chemistry*. 2016 Nov 10;123:236-55.
5. Bhagdev K, Sarkar S. Benzothiazole: As an Antidiabetic Agent. *Annals of the Romanian Society for Cell Biology*. 2021 Jul 10:20269-85.
6. Agarwal S, Gandhi D, Kalal P. Benzothiazole: a versatile and multitargeted pharmacophore in the field of medicinal chemistry. *Letters in Organic Chemistry*. 2017 Dec 1;14(10):729-42.
7. Patel NB, Khan IH, Pannecouque C, De Clercq E. Anti-HIV, antimycobacterial and antimicrobial studies of newly synthesized 1, 2, 4-triazole clubbed benzothiazoles. *Medicinal Chemistry Research*. 2013 Mar;22(3):1320-9.
8. Montalvão S, Leino TO, Kiuru PS, Lillsunde KE, Yli-Kauhaluoma J, Tammela P. Synthesis and biological evaluation of 2-aminobenzothiazole and benzimidazole analogs based on the clathrodin structure. *Archiv der Pharmazie*. 2016 Feb;349(2):137-49.
9. Manfroni G, Meschini F, Barreca ML, Leyssen P, Samuele A, Iraci N, Sabatini S, Massari S, Maga G, Neyts J, Cecchetti V. Pyridobenzothiazole derivatives as new chemotype targeting the HCV NS5B polymerase. *Bioorganic & medicinal chemistry*. 2012 Jan 15;20(2):866-76.
10. Girijavallabhan VM, Alvarez C, Bennett F, Chen L, Gavalas S, Huang Y, Kim SH, Kosinski A, Pinto P, Rizvi R, Rossman R. Synthesis and SAR of pyridothiazole substituted pyrimidine derived HCV replication inhibitors. *Bioorganic & medicinal chemistry letters*. 2012 Sep 1;22(17):5652-7.
11. Landolfo S, Gariglio M, Gribaudo G, Lembo D. The human cytomegalovirus. *Pharmacology & therapeutics*. 2003 Jun 1;98(3):269-97.
12. Borthwick AD, Davies DE, Ertl PF, Exall AM, Haley TM, Hart GJ, Jackson DL, Parry NR, Patikis A, Trivedi N, Weingarten GG. Design and synthesis of pyrrolidine-5, 5'-trans-lactams (5-oxo-hexahydropyrrolo [3, 2-b] pyrroles) as novel mechanism-based inhibitors of human cytomegalovirus protease. 4. Antiviral activity and plasma stability. *Journal of medicinal chemistry*. 2003 Oct 9;46(21):4428-49.

13. Abdel-Aziza HA, Abdel-Wahab BF, Badria FA. Stereoselective Synthesis and Antiviral Activity of (1E, 2Z, 3E)-1-(Piperidin-1-yl)-1-(arylhydrazono)-2-[(benzoyl/benzothiazol-2-oyl) hydrazono]-4-(aryl1) but-3-enes. *Archiv der Pharmazie: An International Journal Pharmaceutical and Medicinal Chemistry*. 2010 Mar;343(3):152-9.
14. Martina BE, Koraka P, Osterhaus AD. Dengue virus pathogenesis: an integrated view. *Clinical microbiology reviews*. 2009 Oct;22(4):564-81.
15. Low JG, Ooi EE, Vasudevan SG. Current status of dengue therapeutics research and development. *The Journal of infectious diseases*. 2017 Mar 1;215(suppl_2):S96-102.
16. Tripathi KD. Essentials of medical pharmacology. JP Medical Ltd; 2013 Sep 30.
17. Al-Soud YA, Al-Sa'doni H, Amajaour HA, Al-Masoudib NA. Nitroimidazoles, Part 3. Synthesis and anti-HIV activity of new N-alkyl-4-nitroimidazoles bearing benzothiazole and benzoxazole backbones. *Zeitschrift für Naturforschung B*. 2007 Apr 1;62(4):523-8.
18. Al-Masoudi NA, Jafar NN, Abbas LJ, Baqir SJ, Pannecouque C. Synthesis and anti-HIV activity of new benzimidazole, benzothiazole and carbonylhydrazide derivatives of the anti-inflammatory drug indomethacin. *Zeitschrift für Naturforschung B*. 2011 Sep 1;66(9):953-60.
19. Kumar M, Chung SM, Enkhtaivan G, Patel RV, Shin HS, Mistry BM. Molecular Docking Studies and Biological Evaluation of Berberine–Benzothiazole Derivatives as an Anti-Influenza Agent via Blocking of Neuraminidase. *International journal of molecular sciences*. 2021 Jan;22(5):2368.
20. De Clercq E, Descamps J, Verhelst G, Walker RT, Jones AS, Torrence PF, Shugar D. Comparative efficacy of antiherpes drugs against different strains of herpes simplex virus. *Journal of Infectious Diseases*. 1980 May 1;141(5):563-74.
21. Field AK, Davies ME, De Witt CM, Perry HC, Schofield TL, Karkas JD, Germershausen J, Wagner AF, Cantone CL, MacCoss M, Tolman RL. Efficacy of 2'-nor-cyclicGMP in treatment of experimental herpes virus infections. *Antiviral research*. 1986 Oct 1;6(6):329-41.
22. Boyd MR, Bacon TH, Sutton DA, Cole MA. Antiherpesvirus activity of 9-(4-hydroxy-3-hydroxy-methylbut-1-yl) guanine (BRL 39123) in cell culture. *Antimicrobial agents and chemotherapy*. 1987 Aug;31(8):1238-42.
23. Amtmann E, Müller-Decker K, Hoss A, Schalasta G, Doppler C, Sauer G. Synergistic antiviral effect of xanthates and ionic detergents. *Biochemical pharmacology*. 1987 May 1;36(9):1545-9.
24. Collins P, Bauer DJ. Relative potencies of anti-herpes compounds. *Annals of the New York Academy of Sciences*. 1977 Mar 1;284:49-59.
25. Färber I, Klinger C, Wutzler P, Thiel KD, Reefschräger J, Herrmann G. Effect of (E)-5-(2-bromovinyl)-and 5-vinyl-1-beta-D-arabinofuranosyluracil on Epstein-Barr virus antigen expression in P3HR-1 cells: comparison with acyclovir. *Acta virologica*. 1987 Jan 1;31(1):13-8.
26. Hutt-Fletcher LM, Balachandran N, LeBlanc PA. Modification of Epstein-Barr virus replication by tunicamycin. *Journal of virology*. 1986 Jan;57(1):117-23.
27. Mitsuya H, Broder S. Inhibition of the in vitro infectivity and cytopathic effect of human T-lymphotrophic virus type III/lymphadenopathy-associated virus (HTLV-III/LAV) by 2', 3'-dideoxynucleosides. *Proceedings of the National Academy of Sciences*. 1986 Mar 1;83(6):1911-5.
28. Lin JC, DeClercq E, Pagano JS. Novel acyclic adenosine analogs inhibit Epstein-Barr virus replication. *Antimicrobial agents and chemotherapy*. 1987 Sep;31(9):1431-3.
29. Lin JC, Smith MC, Cheng YC, Pagano JS. Epstein-Barr virus: inhibition of replication by three new drugs. *Science*. 1983 Aug 5;221(4610):578-9.
30. Lin JC, Smith MC, Choi EI, De Clercq E, Verbruggen A, Pagano JS. Effect of (E)-5-(2-bromovinyl)-2'-deoxyuridine on replication of Epstein-Barr virus in human lymphoblastoid cell lines. *Antiviral research*. 1985 Jan 1:121-6.