Potential of diterpene compounds as antivirals, a review

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ABSTRACT

Viruses cause widely transmitted diseases resulting in pandemic conditions. Currently, the world is being hit by the Covid-19 pandemic caused by the SAR-CoV-2 infection. Countries in the world are competing to develop antivirals to overcome this problem. Diterpene compounds derived from natural ingredients (plants, corals, algae, fungi, sponges) and synthesized products have potential as antivirals. This article summarizes the different types of diterpenes such as daphnane, tiglilane, kaurane, abietane, pimarane, labdane, dollabelane, jatrophane, dolastane, prenylated guaiane, tonantzitlolone, casbane, have antivirals activity such as targeting HIV, Coxsackie virus, herpes virus, hepatitis virus, influenza virus, Chikungunya virus, Zika virus, dengue virus, and SARS-CoV. Some compounds such as andrographolide and its derivatives show promising activity in inhibiting the influenza virus. Additionally, compounds such as pineolidic acid, forskolin, sugiol, and many other diterpene compounds showed anti-SAR-CoV activity. The diterpene compound class's high antivirals potential does not rule out the possibility that these compounds can also act as anti-SAR-CoV-2 drugs in the future.

1. Introduction

Many of the infectious diseases epidemics which becomes a pandemic, which occur due to viral infections. Throughout history, there have been many disease outbreaks caused by viral infections, including Spanish Flu Pandemic (1918–1920), Smallpox (1972), HIV epidemic (1981), SARS (2003), H1N1 Pandemic (2009), Ebola Virus (2014–2016), Zika Virus (2015–2016) (Huremović, 2019). Moreover, in December 2019 there were new cases of pneumonia, and the first reported cases in China were caused by infection with a new virus, novel coronavirus (SARS-CoV-2) known as novel coronavirus disease 2019 (COVID-19) by WHO (Lai et al., 2020). In general, viruses are intracellular parasites containing single-stranded or double-stranded, linear or circular RNA or DNA genomes (Gelderblom, 1996; Gohar et al., 2021).

Pandemics have a direct impact on health and can also adversely affect economic, social, and political stability. The pandemic has infected millions of people and caused thousands of deaths. Travel is strictly restricted, the closure of schools, markets, and specific sectors that will indirectly affect economic, social, security growth, etc (Qiu et al., 2017). The problem of a pandemic is a global problem, and every country is racing to overcome it. The development of antivirals is one of the efforts to accelerate recovery from normal conditions. One of the antiviral compounds that have been widely observed is the terpene group which can be obtained from natural or synthetic materials.

Diterpenes are the product of the mevalonic acid biosynthesis pathway (Singh and Sharma, 2015). Some diterpenes indicated to have antivirals activity, such as kirkinine, excoecariatoxin (anti-HIV) (Olivon et al., 2015), jiadifenoic acids JP (anti-Coxsackie virus) (Zhang et al., 2014), briaexcavatolide U, briaexcavatin L (anti-HCMV) (Yeh et al., 2012), genkwamine P, laurifolioside A (anti-HBV) (Li et al., 2018), linearol, isosidol (anti-HPIV-2) (Kilic et al., 2020), debromoaplysiatoxin (anti-CHIKV) (Gupta et al., 2014) and others. Additionally, several...
Figure 1. Daphnane diterpen as antivirals.
subclasses of diterpene compounds have potential antiviral activity, such as daphnane (Figure 1), tiglane (Figure 2), kaurene (Figure 3), abietane (Figure 4), pimarane (Figure 5), labdane (Figure 6), dolabellane (Figure 7), jatrophan (Figure 8), tonantzitlolone (Figure 9), and miscellaneous (Figures 10 and 11), which are the main focus of this article. The purpose of this review article is to examine the terpenes compounds that have the potential in the development of antivirals to overcome the current pandemic.

2. Type of diterpenes

Diterpenes are chemical compounds consisting of two isoprene units. These compounds are found in animals, plants, fungi, algae, coral, and others. Several types of diterpene compounds such as daphnane, tigliane, kaurene, abietane, pimarane, labdane, dolabellane, jatrophan, dolastane, prenylated guaiane diterpene, tonantzitlolone, and miscellaneous. Diterpene compounds have pharmacological activities such as antivirals. Daphnane diterpene (1–20) is one type of diterpene that generally in the form of a 5/7/6-tricyclic ring equipped with a polyhydroxyl group at positions C3, C4, C5, C9, C13, C14, or C20. Additionally, several diterpene daphnane groups have orthoester groups that bind to C9, C13, and C14 (Jin et al., 2019). Daphnane diterpenes such as acutilobins A, wikstchalide W, Trigocherrins A, and others. This type has Antivirals activity, such as anti-HIV (Yan et al., 2018), and anti-HPIV-2 (Dang et al., 2015).

Abietane diterpene (52–75) is a tricyclic diterpene consisting of about 20 carbons. Compounds belonging to this type include rosmanol, jiadijefenoic acid C, sugiol, and others. Antivirals of this type, among others, as anti-HIV (Pari\textsuperscript{C19}s et al., 1993), anti-Coxsackie (Gonz\textsuperscript{C19}alez and Zaragoza, 2014), anti-ZIKV (Sousa et al., 2020), and anti-DENV (Sousa et al., 2020), and anti-SAR-CoV (Wen et al., 2007). Pimarane diterpene (76–81) is a type of tricyclic diterpene. Pimarane diterpene-type compounds include jiadijefenoic acids L, 4-epi-isopiric acid, forskolin, and others. The reported antivirals for this type are anti-Coxsackie (Zhang et al., 2013) and anti-SAR-CoV (Wen et al., 2007).

Labdane diterpene (82–116) is a bicyclic diterpene. This compound is composed of 4 isoprene units. The basic structure consists of two parts, namely two cyclohexene rings (C1–C10) which are joined to form a decalin system and five-carbon chains (C11–C16) bonded to C-9. There are five methyl branches attached to C4 (for C18 and C19), C-8 (for C17), C10 (for C20), and C13 (for C16) respectively (Tran et al., 2017). The
types of labdane diterpenes include andrographolide, forsythiyanins A, pineolodic acid, and others. This type of antivirals activity is like anti-HIV (Niranjan Reddy et al., 2005), anti-HBV (Chen et al., 2014), anti-DENV (Panraksa et al., 2017), and anti-influenza (Chen et al., 2009).

Dollabelane diterpene (117–131) is one of the diterpenes whose basic structure complies with the 5/11-bicyclic system. This type has anti-HIV (Pardo-Vargas et al., 2014) and anti-HSV (Ogawa et al., 2018). Jatropane diterpene (132–137) is a bicyclic pentadecane diterpene. The basic structure of the diterpene jatrophane consists of four isoprene units forming a ring with a 5/12-bicyclic system. The structure is usually equipped with several substituents such as acyl groups, acetyl, ptopanoyl, butanoyl, isobutyryl, benzoyl, and others (Zhao et al., 2020). Jatrophane equipped with several substituents such as acyl groups, acetyl, ptopanoyl, butanoyl, isobutyryl, benzoyl, and others (Zhao et al., 2020). Jatropane diterpenes had reported having antiviral activity such as anti-HBV (Li et al., 2016). Dolastane diterpene (138–142) is a diterpene composed of a 5/7/6-tricyclic ring system (Tuckett et al., 1998). This type of antivirals activity is anti-HIV (Vallim et al., 2010), anti-HHV, anti-BoHV-5 (Pinto et al., 2019), anti-CHIKV, and anti-ZIKV (Cirne-Santos et al., 2020).

Prenylated guaiane diterpene (143–144) has been reported to be anti-HSV (Kashman et al., 1987). Tonantizitoloine diterpene (145–147) can act as an anti-CHIKV (Olivon et al., 2015). Casbane diterpene (148–149) has anti-HCMV activity (Wang et al., 2013a,b). Miscellaneous diterpene (150–209) is the most common type of diterpene reported as antivirals in this report. This type is reported to be anti-HIV (Pereira et al., 2004), anti-Coxsackie (Gu et al., 2014), anti-HSV (Gonzalez et al., 2010), anti-influenza virus (Dang et al., 2015), anti-HCMV (Wang et al., 2013a,b), and anti-SAR-CoV (Wen et al., 2007) (Table S1).

3. Human Immunodeficiency Virus (HIV)

Human Immunodeficiency Virus (HIV) is a disease caused by a virus that affects the body’s immune system which can lead to AIDS disease. It is recorded that until 2019 there were 75.7 million people infected with HIV from the start of the epidemic with a death rate of 32.7 million people with AIDS (HIV/AIDS, 2020). HIV belongs to the Retroviridae family and the genus Lentivirus, shaped like a ball with a diameter of 100 nm. The virus has two copies of single strain RNA and can transcribe (Lefkowitz et al., 2018). There are two types of Human Immunodeficiency Virus (HIV): Human Immunodeficiency Virus types 1 and 2 (HIV-1 and HIV-2). HIV-1 is known to spread throughout the world, while HIV-2 is spread in West Africa and India (Luzuriaga, 2018). HIV is cytotoxic to CD4+ cells. Infection of CD4+ cells causes a decrease in CD4 cell counts in HIV patients. HIV can be transmitted through sexual contact, blood transfusion, maternal and infant contact, after or before delivery (Levy and Castelli, 2019). HIV enters the host cell through the interaction between the virus’s glycoprotein envelope of the receptor cells on the host cell. The viral RNA genome is inserted into the host cell through membrane fusion. The viral RNA genome undergoes a reverse transcription process with the help of the reverse transcriptase enzyme (Giese et al., 2016). Several diterpene compounds have been indicated to have anti-HIV activity both from the natural product and synthetic.

Daphnane diterpenes such as acutilobins A–G (1–7), daphnethoxin (8), genkwanine VIII (9), kirknine (10), exococcaratxin (11), and 14’-ethylettrahydroxuratoxin (12), isolated from Daphne acutiloba Rehd rods had an anti-HIV-1 activity with an EC50 of less than 2 nM (Huang et al., 2012). The leaves of Stillingia lineata are indicated to contain 12-O-acetylsphorhol-13(2’-methyl)-butyrate (21), 12-O-acetylphorbol-13(2’-methyl)-butyrate (22), 12β-O-[nona-2Z,4E, 6E-trienoyl]-4α-deoxyphorbol-13-butryrate (23), 12-deoxyphorbol-13(2’-methylbutyrate (24), and 12-deoxyphorbol-13-[8’-oxo-hexadeca-2E,4E,6E-trienoate] (25) which functions actively as anti-HIV. Compounds 21–25 can inhibit the outgrowth of HIV-1 and HIV-2 in the strong category, with an EC50 value of less than 1 μM (Olivon et al., 2015).

Anti-HIV-1 and anti-HIV-2 activity through inhibition of viral replication has been indicated from the terpenoid compounds of the Euphorbiaceae species. Phorbol ester derivative compounds, phorbol-12,13-diacetate (30), phorbol-12,13-dodecanoate (26), phorbol-12,13-dodecanoate (27), phorbol-12,13-tetradecanoate (29), phorbol-12,13-diacetate (30), phorbol-12,13-dibutyrate (31), phorbol-12,13-dihexanoate (32), phorbol-12,13-didecanoate (33), 12-O-tetradecanoylphorbol-13-acetate (34), 12-O-tiglylphorbol-13-dodecanoate (35), 12-O-(N-methylanthranilate)-phorbol-13-acetate (36), 12,13-O, O’-dinoanoylphorbol-20-
homovanillate (37), 12-O-phenylacetyl-13-O-acetylphorbol-20-homovanillate (38), phorbol-12,13,20-triacetate (39), and 12-O-tetradecanoyl-20-oxo-20-deoxyphorbol-13-acetate (40) has anti-HIV-1 activity with IC\(_{50}\) values ranging from 0.9 nM - 27.4 \(\mu\)M, with compound 29 having the best activity. 12-deoxy phorbol derivative, 12-deoxyphorbol-13-acetate (prostratin) (41), 13-O-isobutyryl-12-deoxyphorbol-20-acetate (42), and 13-O-phenylacetyl-12-deoxyphorbol-20-acetate (43), also acts as anti-HIV-1 with an IC\(_{50}\) value of 0.2 \(\mu\)M–1.9 \(\mu\)M. Ingerol ester derivative compounds, ingenol-3-mebutate (ingenol-3-angelate) (202), and ingenol-3,20-dibenzoate (203) able to inhibit HIV-1 activity with IC\(_{50}\) values 17 nM and 27 nM. Apart from being active as anti-HIV-1, compounds 27–44 and 202–203 also have anti-HIV-2 activity. Compounds 29, 31–38, 43, 202, and 203 have HIV-2 inhibitory activity with IC\(_{50}\) values ranging from 0.2 nM - 80 nM, with the best inhibitory activity shown by compounds 23 and 35 (IC\(_{50}\) 0.2 nM). The anti-HIV mechanism is thought to be related to the activation of kinase C isoenzymes (PKC) (Nothias-Scaglia et al., 2015). Furthermore, anti-PKC beta antibodies inhibited phorbol-12-myristate acetate (44) induced activation of NF-\(\kappa\)B and HIV-1 (Kim et al., 1996).

Diterpenes from sponge Hyattella intestinalis, spongiatriol (152), and isospongiatriol (153) were significantly able to inhibit HIV growth with IC\(_{50}\) values less than 20 \(\mu\)g/ml. The two diterpenes act as inhibitors against the activation of NF-\(\kappa\)B (Nuclear Factor kappa light chain enhancer of activated B cells) (Ahmadi et al., 2017). NF-\(\kappa\)B is one of the transcription factors that is activated by a viral infection. NF-\(\kappa\)B is required to express IFNB (Interferon Beta), induced by viruses (Balachandran and Beg, 2011). Ent-3S-hydroxy-atis-16(17)-ene-1,14-dione (154) isolated from Homalanthus nutans, ent-15-oxo-kaur-16-en-19-oic acid (45) and ent-15-oxo-kaur-16-en-19-oic acid methyl ester (46)
isolated from *Chrysobalanus icaco* has anti-HIV activity with EC\textsubscript{50} value 19, 1.58 and 50 \(\mu\)M respectively (Gustafson et al., 1991). Euphorneroid D (150) and *ent*-3-oxoatian-16\(\alpha\),17-acetonide (151) from the bark of *Euphorbia neriifolia* showed anti-HIV-1 activity with EC\textsubscript{50} 34 \(\mu\)M and 24 \(\mu\)M (Yan et al., 2018).

*Tripterygium wilfordii* root contains tripterifordin (47) also has potential as an anti-HIV-1 (Chen et al., 1992, 1995). 16\(\beta\),17-Dihydroxy-ent-kauran-19-oic acid (48) from *Annona squamosal* (Wu et al., 1996) and methyl-16\(\alpha\)-hydro-19-al-ent-kauran-17-oate (49) from *Annona glabra* (Chang et al., 1998) able to inhibit HIV-1 replication with EC\textsubscript{50} value 2.38 \(\mu\)M and 15 \(\mu\)M, respectively, in H9 lymphocytes. Carnosalic acid (52) from *Rosmarinus officinalis* can inhibit HIV-1 protease by 90%. While the derivatives, rosmanol (53), 7-O-methylrosmanol (54), and 7-O-ethylrosmanol (55) also have anti-HIV-1 activity. The existence of a benzilic group CH\(_2\) and a carboxylic group performs in the inhibiting of HIV-1 protease from carnosic acid (Pari\textsubscript{1}/C20\textsubscript{s} et al., 1993).

Figure 6. Labdane diterpene as antivirals.
The dolabellane diterpenes of the octocoral Caribbean *Eunicella laciniata* are 13-keto-1(R),11(S)-dolabella-3(E),7(E),12(18)-triene (117), and β-Araneosene (118) exhibits HIV-1 inhibitory activity. In addition, the derivatives of β-Araneosene, (1R,11S,3S,4S,7S,8S)13-keto-di-3,4:7,8-epoxy-dolabella-12(18)-ene (119), (1R,3R,4R,7S,11S)13-keto-3,4,7-trihydroxy-dolabella-8(17),12(18)-diene (120) and (1R,3E,7E,11S)-13-keto-16-hydroxy-dolabella-3,7,12(18)-triene (121) showed an impressive increase in anti-HIV-1 activity compared to compound 117 with EC50 value 0.73 μM, 3.9 μM and 0.69 μM (Pardo-Vargas et al., 2014). Additionally, the compound of 8,10,18-trihydroxy-2,6-dolabelladiene (122) from the algae *Dictyota pfaffii* was indicated to have anti-HIV activity. This compound acts as a non-competitive inhibitor of the HIV-1 reverse transcriptase enzyme so that it can interfere with HIV-1 replication (Cirne-Santos et al., 2006, 2008). Reverse transcriptase is a multifunctional enzyme such as RNA-dependent DNA polymerase, DNA-dependent DNA polymerase, and RNase H activities (Pomerantz and Horn, 2003).

Andrographolide (82) can reduce the amount of p24 antigen on MT2 cells so that it has the potential to be an anti-HIV drug (Niranjan Reddy et al., 2005). This compound can also inhibit HIV-induced cell-cycle dysregulation, this effect in a significant increase in the concentration of CD4+ lymphocytes so that it can reduce replication. HIV (Calabrese et al., 2006). Dehydroandrographolide succinic acid monoester (84) compound has HIV-1 and HIV-2 inhibitory activity by inhibiting HIV-induced cell fusion (Chang et al., 1991).

The SJ23B (187) compound isolated from *Euphorbia hyberna* is classified as a jatrophane diterpenes capable of inhibiting HIV-1 infection with an IC50 value of 2 nM. The compounds’ mechanism is to induce the latent reactivation process of HIV-1 by activating NF-κB and down-regulating the expression of HIV-1 receptors such as CD4, CXCR4, and CCR5, thereby preventing infection of other CD4+ cells (Bedoya et al., 2009). Two diterpenes, (6R)-6-hydroxydichotoma-3,14-diene-1,17-dial (204) and (6R)-6-acetoxidichotoma-3,14-diene-1,17-dial (205) from *Dictyota menstrualis* have anti-HIV-1 activity by inhibiting the replication of the virus. The diterpene compound can inhibit the viral replication process by acting as an inhibitor of the HIV-1 reverse transcriptase enzyme (Pereira et al., 2004).

4. Coxsackievirus

Coxsackievirus is part of the enterovirus genus of the Picornaviridae family. This virus is 30 nm in size and has single-stranded RNA. Furthermore, there are two types of Coxsackievirus, including Coxsackievirus A and B (Oberste, 2008). This virus can be transmitted through respiratory droplets, blisters, or feces (Ang et al., 2009). This disease causes hand, foot, and mouth disease (HFMD) since 1997 (Lam et al., 1998). The Study of compounds diterpenes as anti-Coxsackie viruses have long been used.

The terpenic abiatane compound from *Illicium jiadifengpi* has anti-Coxsackie activity. Jiadifenolic acids J-K (56–57), jiadifenolic acids L-N...
(76–78), and jiadifenoic acids O–P (155–156) have anti-Coxsackie virus type B3 activity. The IC_{50} value of compounds 77, 78, and 156 against Coxsackievirus inhibition were less than 20 μmol/mL (Zhang et al., 2014). α-hydroxycallitrisic acid (58), 4-epi-isomapiric acid (79), and isopimara-7,15-dien-19-ol (80) significantly showed antivirals activity against Coxsackievirus types B2, B3, B4, and B6 with an IC_{50} value of less than 25 μmol/mL (Zhang et al., 2013). Abietane diterpenes (+)-jiadife-noic acid C (59) which is a semisynthetic product of callitrisic acid has anti-Coxsackie B virus (González/C19alez and Zaragoza/C19a, 2014). The root of Illicium majus contains 15S-majusanic acid E (60) and 15R-majusanic acid E (61). These compounds had anti-Coxsackie B3 virus activity with IC_{50} values of 17.4 μg/mL and 12.8 μg/mL (Wang et al., 2013a,b). The terpilisabolane compound from Claoxylon polot, namely claoxylones A-I (188–196) showed anti-Coxsackie B3 virus activity with an IC_{50} value of 6.0–33.3 μM (Gu et al., 2014).

5. Herpes virus

Herpes Simplex Virus (HSV) belongs to the Herpesviruses family and the genus Simplexviruses (Knipe and Whitley, 2020). There are two types of herpes simplex virus (HSV), herpes simplex virus types 1 and 2 (HSV-1 and HSV-2). HSV-1 infects the ororalabial area, whereas HSV-2 targets the genital area (Widener and Whitley, 2014). HSV-1 and HSV-2 are frequent, long-term infections, often asymptomatic (Tronstein et al., 2011). It is estimated that in 2016 there were 491 million cases of HSV-2 that infected people aged 15–49 years. HSV-1, it is estimates that there are 192 million cases (James et al., 2020). Several articles have observed anti-HSV activity in several terpenes.

The scopadurosane (157) derivative compound, the compound 158–160 has anti-HSV-2 activity. The presence of a hydroxy group (-OH) on C-13 in compound 158–160 a affects the compound's anti-HSV-2 strength (González et al., 2010). Dolabellane diterpenes, (1R,3R,4R,7R,8R,11S)-di-3,4:7,8-epoxy-13-keto-dolabell-12(18)-ene (123) from Eunicea laciniata and (1R,3E,4R,7R,8R,11S)-di-7,8-epoxy-13-keto-dolabell-3,12(18)-diene (124) from Eunicea sperula has anti-HSV-1 activity with inhibition values of 42.6% and 73.7% (Amaya-Garcia et al., 2017). Nigella damascene seeds contain dolabellane diterpenes, damasterpenes I-III (129–131) and damasterpenes V-VIII (125–128). Compounds 125, 126, and 130 showed HSV-1 inhibitory activity at a concentration of 10 μM with 20–35% inhibition. Meanwhile, compounds 127, 128, 129, and 131 showed less than 20% (Ogawa et al., 2018).

Two dolastane diterpenes from the brown algae Canistrocarpus cervicornis, 4-hydroxy-9,14-dihydroxydolasta-1(15),7-diene (138) and 4,7,14-trihydroxydolasta-1(15),8-diene (139) can inhibit HSV-1 infection with 90% and 99% inhibition. Both of these compounds can become anti-HSV-1 agents because their inhibition values are similar to acyclovir (inhibition = 99%) (Vallim et al., 2010). Anti-herpesvirus activity in two dolastane diterpenes (4R,7R,14S)-4α,7α-diacetoxy-14-hydroxydolast-1(15),8-diene (140) and (4R,9S,14S)-4α-acetoxy-9β,14α-dihydroxydolast-1(15),7-diene (141) from Canistrocarpus cervicornis. Compound 140 was reported to have much better inhibitory activity on human alphaherpesvirus-1 (HHV) than compound 141 with...
EC_{50} values 6.25 \mu M and 120 \mu M. Additionally, these compounds can also inhibit Bovine alphaherpesvirus 5 (BoHV-5) (Pinto et al., 2019). Dolastane diterpene compounds can inhibit the HSV replication process by inhibiting DNA polymerase activity (Vallim et al., 2010). Meanwhile, two prenylated guaiane compounds from Epipolasis reiswiq, reiswigins A and B (143–144) were also indicated to have anti-HSV-1 activity (Kashman et al., 1987).

Briareum excavatum has been reported to contain briacavatolides A–C (182–184), briacavatolide U (185), and briacavatin L (186). Anti-HCMV evaluation showed that the briacavatolide C (184) compound had the greatest inhibitory activity with an IC_{50} value of 18 \mu M (Yeh et al., 2012). Ehrenbergol C and asetil ehrenberoksida B (148–149) from soft coral Sarcophyton ehrenbergi had anti-human cytomegalovirus (HCMV) activity with EC_{50} values 20 and 8 \mu g/mL (Wang et al., 2013a,b). HCMV is beta herpes that causes lifelong infection in humans. This virus is a significant cause of defects in infants and immune disorders. Infants can be infected with this virus through the placenta, labor, and breastfeeding (Landolfo et al., 2003). The gyrosanols A (197) and gyrosanols B (198) from soft coral Sinularia gyrosa have anti-HCMV activity with IC_{50} values 2.6 dan 3.7 \mu g/mL. Also, compounds 197 and 198 show

Figure 10. Miscellaneous diterpene as antivirals.
significant anti-inflammatory activity by reducing the COX-2 protein (Cheng et al., 2010).

6. Hepatitis virus

Hepatitis virus is the major cause of inflammatory liver disease. There are five types of hepatitis viruses, including hepatitis A (HAV), B (HBV), C (HCV), D (HDV), and E (HEV) viruses. Hepatitis A (HAV) and hepatitis E (HEV) are transmitted orally or through feces. Both viruses cause acute illness. Hepatitis B (HBV) and hepatitis C (HCV) are transmitted parenterally and cause chronic hepatitis. Meanwhile, hepatitis D (HDV) is pathogenic when combined with HBV (Lefkowitch, 2021). All types of hepatitis belong to different families. HBA belongs to the Picornaviridae family, HBV from the Hepadnaviridae family, HCV from the Flaviviridae family, and HEV from the Hepeviridae family (Ryu, 2016). Hepatitis virus has caused 1.34 million deaths in 2015, of the total deaths caused by HBV (66%), HCV (30%), HAV (0.8%), and HEV (3.3%) (WHO, 2017).

Wikstroemia Chamaedaphne has been reported to contain diterpene compounds, wikstchalide W (13), genkwamine F (14), aurifolioside A (132), 2-epi-laurifolioside A (133), laurifolioside B (134), 2-epi-laurifolioside B (135), laurifolioside (136), and 2-epi-laurifolioside (137). Compounds (14) and (132) have anti-HBV activity by inhibiting HBsAg surface antecedents with IC50 values of 46.5 and 88.3 μg/mL. While compounds 13, 133–137 were able to inhibit HBV replication in the range of 2%–33% (Li et al., 2018).

Labdane diterpene, andrographolide (82) can inhibit HCV replication by regulating heme oxygenase-1 expression, thereby increasing levels of biliverdin metabolites. This promotes the antivirals response of IFN and inhibits NS3/4A protease activity. Besides, andrographolide can activate the phosphorylation of p38 MAPK, which stimulates the expression of heme oxygenase-1 mediated by nuclear factor erythroid 2 (Nrf2) (Lee et al., 2014). NS3/4A protease is an enzyme responsible for the selective cleavage of polyproteins into individual viral proteins. This protease plays an essential role in HCV replication, one of the three HCV drug targets (Meewan et al., 2019).

Andrographolide (82) and dehydroandrographolide (83) from Andrographis paniculata can inhibit HBV DNA replication with IC50 value 22.6 μM and 54.1 μM. A total of 48 derivatives of the two compounds were evaluated for anti-HBV activity. There were 21 derivatives of dehydroandrographolide and andrographolide (95–115) with IC50 values less than 50 μM with anti-HBV mechanisms by inhibiting viral DNA replication (Chen et al., 2014). Compounds 95, 99, and 112 had anti-HBV activity by inhibiting HBsAg secretion with IC50 value less than 50 μM (Chen et al., 2014). HBsAg (Hepatitis B surface antigen), is a hepatitis B virus surface antigen consisting of lipids and proteins on the surface of the virion and is produced excessively during the virus life cycle (Blumberg and Alter, 1965; Howard, 1986). The mechanism for inhibiting HBsAg secretion is shown by compound 95 and 100 with IC50 values less than 50 μM (Chen et al., 2014). Hepatitis B e antigen (HBeAg) is a product of the nucleocapsid (core) HBV gene secreted in serum. HBeAg is a marker of viral replication associated with liver infectivity and damage (Karayiannis et al., 1985).

7. Influenza virus

Influenza is an RNA virus that includes in the Orthomyxoviridae family. Influenza viruses consist of three types include influenza A, B, and C. This virus has three glycoprotein envelopes: hemagglutinin (HA), neuraminidase (NA), and matrix protein (M1 and M2). The influenza virus has eight RNA strands that allow the formation of new virus subtypes. For example, the H1N1 influenza virus has a single-stranded RNA genome, negative sense, and ribonucleoproteins of different sizes (Zhang and Lamb, 1996).

Antiviral activity against Human Parainfluenza Virus Type 2 (HPIV-2) was also investigated in the enk-kaurene diterpenes. Linearol (50) and isosidol (51) compounds obtained from the aerial part of Sideritis Lycisti have anti-HPIV-2 activity with an antiviral index value (CD50/ED50) of less than 2 μM (Kilic et al., 2020). Phenolic derivatives of diterpenes (+)-acids podocarpic (161) and (+)-totarol (181) have anti-influenza A virus activity. Evaluation of (+)-podocarpic acid derivatives (161–181) can inhibit the influenza virus. Methyl podocarpate (162) and its O-acetyl derivative (177) can increase the anti-influenza virus activity of podocarpic acid (161) by 100 times stronger. Meanwhile, benzyl ester (163) can increase anti-influenza virus activity 10 times stronger. Totarol

Figure 11. Miscellaneous diterpene as antivirals (continuation).
has 5 times stronger activity to inhibit influenza virus (PR8) compared to podocarpic acid. The totoral was able to inhibit about 12 times stronger VR1679 virus (H3N2) than compound 170. However, compound 170 inhibited PR8 virus (H1N1) 15 times stronger than totoral. Compound 164 has anti-influenza virus (PR8) activity through hemolysis inhibition mediated by hemaggiltamin. Hemaggiltamin protein is essential for virus entry (Dang et al., 2015).

Andrographolide (82), 14-dehydroxyandrographolide-12-sulfonic acid sodium salt (85), and 14-a-lipoyl andrographolide (86) can inhibit various kinds of influenza viruses such as H5N1, H1N1, and H9N2 (Chen et al., 2009). Andrographolide can block the interaction pathway between the retinoic acid-inducible gene-1 (RIG-1) receptors induced by the influenza virus. In this process, there was an improvement in cell death caused by influenza virus infection (Yu et al., 2014). The 14-a-lipoyl andrographolide (86) was able to inhibit virus adsorption to red blood cells with a minimum inhibitory concentration (MIC) of 5.3 mM (H1N1), 7.1 mM (H9N2), and 16.8 mM (H5N1). This shows that the compound can interfere with viral hemagglutinin so that it can block binding to cellular receptors (Chen et al., 2009).

Some labdane diterpene isolated from the fruit of Forsythia suspensa have been indicated to have anti-influenza activity type H1N1 virus. 3β-hydroxy-8(17),13E-labadien-15-α-oic acid (87), forsyihyisanins A-B (88–89), 19-hydroxy-ent-labda-8(17),13E-dien-15-α-oic acid (90), ent-linda-8(17),13E-dien-15,19-dioic acid (91), ent-linda-8(17),13Z-dien-15,19-dioic acid (92), manio-labda-8(20),13-dien-15,18-dioic acid (93), and 18-hydroxy-7-oxolabda-8(9),13E-dien-15-α-oic acid (94), all of these compounds have IC₅₀ values of less than 30 μM (Zhao et al., 2020).

Rhododromin B (142) from Rhododendron molle fruits showed inhibitory activity of Influenza A95/359 virus with an IC₅₀ value of 19.24 μM (Li et al., 2016). *Trichoderma atroviride* containing wickerols A and B (200–201) had anti-influenza virus H1N1 strain with IC₅₀ values 5.0 μg/ml and 0.07 μg/ml. Both compounds can inhibit DNA polymerase activity in viruses so that the viral replication process is inhibited (Yamamoto et al., 2012).

8. Chikungunya virus (CHIKV)

Chikungunya virus (CHIKV) is a virus, including in the Flaviviridae family. *Aedes aegypti* mosquito spreads the virus. The incubation period for CHIKV is relatively short, 2–4 days (Halstead, 2018). CHIKV disease is characterized by high fever, pain in one or several joints, and a rash on the face (De Ranitz et al., 1965). CHIKV infection is widespread in various countries in Asia (Jamal and Sam, 2018), Europe (Zanoni et al., 2017), America (Vasconcelos et al., 2018), and Africa (Diallo et al., 2018).

Several diterpenes have indicated antiviral activity against CHIKV. Daphnane diterpenes from *Trigonostemon cherrieri*, trigocherrins A-B (15–16), trigocherrin F (17), and trigocherrilides A-C (18–20) can inhibit CHIKV virus replication better than chloroquine (Allard et al., 2012). Daphnane diterpenes can reduce viral protein gene expression so that CHIKV RNA production decreases. The decreased production of CHIKV RNA was also caused by blocking the interaction between viral DNA and the reverse transcriptase enzyme so that the viral replication process was inhibited (Kaur et al., 2013). Meanwhile, tonantlzitilobone, tonantlzitiltin C, and tonantlzitiltin F (145–147) from *Stilingia lineata* leaves can inhibit CHIKV by EC₅₀ value less than 25 μM (Oliven et al., 2015).

The Anti-CHIKV activity of several terpenes are from the marine cyanobacterium *Trichodesmium erythraeum*, debrumosalphatosiytoxin (207), anhydrodebrumosalphatosiytoxin (208), and 3-methoxydebrumosaplysoiytoxin (209). Compound 207 can inhibit CHIKV infection better than compounds 208 and 209. These three compounds inhibit the viral replication process after the virus enters the host cell by inhibiting viral DNA polymerase activity, which has a necessary role in the viral replication process (Gupta et al., 2014). Dolastane (141) from *Canistrocarpus cervicornis* had an anti-CHIKV activity with an EC₅₀ value of 1.28 μM (Cirne-Santos et al., 2020).

9. Zika virus (ZIKV)

Zika virus (ZIKV) is a virus from the Flaviviridae family that can be transmitted through *Aedes biete, Aegypti* and *Aedes albopictus*. The disease caused by ZIKV infection is almost the same as the dengue virus, West Nile virus, Japanese encephalitis virus, and yellow fever virus but neurological complications can lead to Guillain-Barré syndrome and congenital malformations that cause microcephaly (Pflourde and Bloch, 2016; Noorbakhsh et al., 2019). Zika infection is sometimes asymptomatic, but some cause clinical symptoms non-specifics such as headache, low-grade fever, arthralgia, rash, and conjunctivitis (Le, 2016). Globally, it is estimated that there were 3–4 million cases of Zika in 2016 (WHO, 2016).

Semi-synthetic derivatives of abieten diterpene, 18-aminoferuginol (71), 12-hydroxy-N,N-phthalyldehydroabietylamine (72), 12-acetoxy-N,N-tetralcholorophthalyldehydroabietyl amine (73), 12-hydroxy-N-tosyl-dehydroabietylamine (74), 18-oxoferuginol (62), and 12-nitro-N-benzoyl-dehydroabietylamine (75) are potential anti-ZIKV agents. The best ZIKV inhibitory activity was compound 75 with EC₅₀ 0.67 μM, while the anti-ZIKV activity of compound 62, 71–74 with EC₅₀ 2.60–18.57 μM (Sousa et al., 2020). Dolastane (141) from *Canistrocarpus cervicornis* had an anti-ZIKV activity with an EC₅₀ value of 0.75 μM (Cirne-Santos et al., 2020).

10. Dengue virus (DENV)

Dengue virus (DENV) is a virus, including in the flavivirus genus and the Flaviviridae family. DENV can cause dengue to dengue shock syndrome (Murugesan and Manoharan, 2020). This virus is spread in tropical areas. There are five types of dengue viruses, including DENV-1, DENV-2, DENV-3, DENV-4, and DENV-5 (Gubler and John, 2014). It is estimated that around 390 million dengue fever cases per year (Bhatt et al., 2013). There is an increase in dengue fever cases annually from 2000 (505,430 cases) to 2019 (4.2 million cases), with deaths increasing from 960 cases (2000) to 4032 cases (2015) (WHO, 2014).

Andrographolide (82) has anti-DENV-2 activity with EC₅₀ 21.30 μM (HepG2 and 22.74 μM (HeLa) (Panraksa et al., 2017). Anti-DENV andrographolide activity is associated with GRP78 and the unfolded protein response (Paemanee et al., 2019). The compound of *Trigonocestemon cherrieri* is trigocherrins A (15), and trigocherrilides A-B (18–19) can inhibit DENV. The IC₅₀ of the three compounds ranged from 3.1 to 16.0 μM (Allard et al., 2012). The anti-DENV activity was also shown by (+)-feruginol (63) and 18-oxoferuginol (62), with inhibition values of 1.4 and 5.0 μM (Sousa et al., 2020).

11. Coronavirus (SARS-CoV)

SARS-CoV (Severe Acute Respiratory Syndrome-Coronavirus) is one of the first infectious diseases discovered in China. This disease is one of the causes of global problems because it spreads in 26 countries on five continents. More than 8,000 SARS cases and 744 death cases have been recorded (Peiris et al., 2003). This virus comes from the Coronaviridae family and has 90 single-stranded RNA genetic material. SARS-CoV attacks the respiratory system. The virus can be transmitted through body fluids (mouth, nose, or eyes). SARS-CoV binds to the ACE2 receptor and introduces its genetic material to host cells by endocytosis. Patients infected with this virus are characterized by a high fever (more than 38 °C), headaches, body aches, and respiratory infections. Symptoms of the disease will appear after 2–14 days after infection (Freeman and Kenez, 2018).

Several diterpenes have the potential to be anti-SARS-CoV. Ferruginol (63), dehydroabietyl-7-one (64), sugiol (65), cryptojaponol (66), [8]-hydroxyabieta-9(11),13-dien-12-one] (67), [7]-hydroxyabietyl-9(11),13-dien-12-one] (68), 6,7-dehydroxyabieta-9(11),13-dien-12-one] (69), 12,13-diacetoxyabieta-6,8,11,13-tetraene (70), pinusolidic acid (116), forskolin (81), cedrene-3α,12-diol (206), and α-cadinol (199) had anti-SARS-CoV activity.
Compounds 63–65, 67, 69, 81, 116 obtained from the wood core of Chamaecyparis obtusa var. formosana, compound 66 obtained from the wood core of Cryptomeria japonica, compound 70, 206, and 199 obtained from the wood core of Juniperus formosana. Compounds 63, 65, 68, and 70 showed significant anti-SARS-CoV activity with EC$_{50}$ values less than 1.5 μM and better than valinomicyn. Meanwhile, compounds 64, 69, 116, 68, and 206 showed moderate activity against SARS-CoV with EC$_{50}$ values between 3.8-7.5 μM. These compounds act as competitive inhibitors against the SARS-CoV 3CL protease by blocking the SARS-CoV 3CL protease activity, thereby disrupting the viral replication cycle. The SARS-CoV 3CL protease is the main target in anti-SARS-CoV treatment. The SARS-CoV 3CL protease has an important role in the virus replication process because it plays a role in viral cell division (Wen et al., 2007).

Inhibition of SARS-CoV by diterpenes at the molecular level through the mechanism of inhibition of the main protease enzyme active site (Figure 12). The targeting of the active site is expected to provide an overview of the inhibition of the main protease 3CL through the interaction between diterpene compounds (ligand) and amino acid residues on the active site of the protein (receptor) (Wen et al., 2007). This interaction is expected to change the conformation of the complex structure (ligand-receptor) so that the protein becomes inactive and stops virus replication. More specifically, the mechanism of inhibition occurs through binding to amino acids on the active site of the main protease as a receptor in the form of hydrogen bonds.

12. Conclusions

Diterpenes are compounds that can be obtained from the natural product or the synthesis process. The source of diterpenes from the natural product can obtain from plants (Daphne acutiloba, Stilingia lineata, Illicium jiadifengyi, Wikstroemia chamaedaphne, etc), porifera (Hyattella intestinalis), coral (Eunicea laciniata, Eunicea laciniata, Briareum excavatum, etc), algae (Dictyota paffii, Canistrocarpus cervicornis, Trichodesmium erythraeum, etc), and fungi (Trichoderma atroviride). Diterpenes antivirals activity can inhibit HIV, Coxsackievirus, HSV, HHV, BoHV-5, HCMV, Hepatitis virus, HPIV, H3N2, H1N1, PR8, H5N1, H9N2, CHIKV, ZIKV, DENV, and SAR-CoV. This class of compounds may be an alternative in developing anti-SARS-CoV-2 drugs to tackle the Coronavirus disease 2019 (Covid-19) pandemic.

Declarations

Author contribution statement

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