MORONIC ACID: A REVIEW

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ABSTRACT

Moronic acid is a pentacyclic triterpenoid made up of olean-18-ene with an oxo group at position 3 and a carboxy group at position 28. It’s made from an oleanane hydride. A few investigations have demonstrated that Moronic acid has a wide range of pharmacological effects such as Antidiabetic activity, Anti-AIDS agents, Chemotherapeutic agents, Virus lytic, Anti-HIV, Cytotoxic activity, Anti-herpes, Antimicrobial activity, Ribosome-loaded mRNAs.

INTRODUCTION

Moronic acid is a naturally occurring triterpene. Moronic acid may be extracted from Rhus javanica, a succulent plant that has traditionally been believed to have medicinal qualities. The chemical has also been extracted using mistletoe (Phoradendron reichenbachianum). Bevirimat, a derivative of the related triterpenoid beta-linolic acid, is being researched as an anti-HIV drug; however, moronic acid has also been shown in vitro to have better antiviral activities than bevirimat. A moronic acid derivative with EC50 values of 0.0085 M against NL4-3, 0.021 M against PR-R (a multiple protease inhibitor resistant strain), and 0.13 M against FHR-2 (a ribavirin-resistant HIV strain) showed potent anti-HIV activity (bevirimat). This derivative, which is also effective against the herpes simplex virus 1, has emerged as a promising new candidate for clinical trials [1, 2].

Chemical structure moronic acid [1]

Synonyms [1]
Moronic acid
6713-27-5
Moronicacid
3-Oxoolean-18-en-28-0ic acid
MLS00056340
(4aS,6aR,6bR,8aR,12aR,14aS)-2,2,6a,6b,9,9,12a,14,14a-hexadecahydro-3H-picene-4a-carboxylic acid

Chemical properties of moronic acid

Appearance: Powder, Formula: C30H46O3, Molecular Weight: 454.7, Type of Compound: Triterpenoids, Storage: Desiccate at -20 °C, Solubility: Soluble in Chloroform, Dichloromethane, Ethyl Acetate, DMSO, Acetone, etc. Generally, Warm the tube to 37 °C and shake it in the ultrasonic bath for a while to increase solubility. The stock solution may be kept for many months at -20 °C. We suggest that you make the solution and utilise it on the same day. If the test schedule demands it, the stock solutions may be produced ahead of time, but they must be sealed and kept below -20 °C. The stock solution may be maintained for many months in most cases. We suggest allowing the vial to come to room temperature for at least an hour before using it.

Source of moronic acid

This product is isolated and purified from the herbs of Rhus chinensis

Pharmacological activities of moronic acid

The pharmacological activities of Moronic acid are briefly discussed are Antidiabetic activity

The anti-diabetic activity of four structurally similar triterpenic acids: ursolic (RE-01), oleanolic (RE-02), moronic (RE-03), and morolic (RE-04) acids, when taken orally. The anti-diabetic effects of these triterpenes (50 mg/kg) on STZ-nicotinamide diabetic rats were evaluated in an acute experiment. In compared to the control group, all drugs exhibited substantial anti-diabetic efficacy (p<0.05). Compounds’ inhibitory efficacy against protein tyrosine phosphatase 1B (PTP-1B) was also tested in vitro. The enzymatic activity was nearly totally suppressed at 50 M. The crystal structure of PTP-1B was used to dock all of the compounds. The triterpenic acids may bind in a binding pocket adjacent to the catalytic site, according to docking studies. The protein-ligand complexes are stabilised by a large hydrogen bond network with the carboxyl group and Van der Waals interactions [3]. The primary components of acetonide extract from Phoradendron reichenbachianum (Loranthaceae), a medicinal plant used in Mexico to treat diabetes, are morolic (1) and moronic (2) acids. The goal of this investigation was to see if compounds 1 and 2 have sub-acute anti-diabetic and antihypertensive effects in a non-insulin-dependent diabetic rat model. Also, an oral glucose tolerance test was used to assess the antihyperglycemic effect on normoglycemic rats. Daily administration of morolic (1) and moronic (2) acids (50 mg/kg) reduced blood glucose levels by 60% from the first to the tenth day following treatment compared to the control group (p<0.05). Furthermore, blood samples from diabetic rats revealed that both substances reduced plasmatic cholesterol (CH0) and triglyceride (TG) concentrations, restoring them to normal levels (p<0.05). Also, compared to the control group, pretreatment with 50 mg/kg of each drug resulted in a substantial antihyperglycemic impact following glucose and sucrose loading (2 g/kg). Compounds 1 and 2 inhibited 11-HSD 1 activity in vitro at 10 M, according to in vitro experiments.
Anti-AIDS agents

Different C-3 conformationally constrained betulinic acid (BA, 1) derivatives were developed and synthesized in order to explore the conformational space of the C-3 pharmacophore in our ongoing investigation of triterpene derivatives as effective anti-HIV medicines. Analogues of 3-0-monomethylsucinyl-betulinic acid (MSB) were also created to better understand the role of the C-3’-dimethyl group in bevirimat (2), a first-in-class HIV maturation inhibitor now undergoing phase Ib clinical trials. In addition, the backbone and C-3 alteration of another triterpene skeleton, moronic acid (MA, 3), were studied in relation to the anti-HIV action of this chemical family. This research helped us better understand the structure-activity relationships (SAR) of triterpene-derived anti-HIV drugs, leading to the design and production of compound 12 (EC50: 0.0006 microM), which had somewhat greater HIV-1 maturation inhibitor efficacy than compound 2 [5].

Seven novel triterpene derivatives were developed, synthesized, and tested for in vitro antiviral activity as part of a continuous structure-activity relationship research of powerful anti-HIV medicines. Moronic acid derivatives 19, 20, and 21 were shown to have considerable action in H9 cells infected with HIV-1. In the MT-4 cell line, compounds 19 and 20 were tested against HIV-1 NLA-4 and treatment resistant strains. The antiviral properties of compounds 19 and 20 were superior to that of the betulinic acid analogue 8 [PA-457], which has completed a Phase IIa clinical study. Compound 20 had EC50 values of 0.0085 microM against NLA-4, 0.021 microM against PR-R (a multiple protease inhibitor resistant strain), and 0.13 microM against PR-RR-2, indicating strong anti-HIV action (an HIV strain resistant to 8). Compound 20 has turned into a fresh lead for modification, and more research into 20-related compounds as clinical trial candidates is needed [2].

From Brazilian propolis, researchers extracted a novel triterpenoid called melliferone (1), three known triterpenoids, moronic acid (2), amuvuwetonic acid (3), and betulinic acid (4), and four known aromatic compounds (5-8) that were evaluated for anti-HIV activity in H9 cells. Moronic acid (2) was modified to create more powerful anti-AIDS drugs after showing substantial anti-HIV activity (EC50 = 0.1 microg/ml, TI = 186) [6].

Chemotherapeutic agents

Pharmaceutical substances have always been abundant in medicinal plants. As a result, the author’s research program’s long-term goals are to find and create novel chemotherapeutic medicines based on plant-derived chemical leads utilising a medicinal chemistry method, which combines chemistry and biology. Sesquiterpene lactones, quassinoids, naphthoquinones, phenylquinolones, dihydroxy-3',7-dimethoxyflavone, were isolated from the cytotoxic MeOH extract obtained from Acridocarpus. On the basis of thorough investigations of the novel compounds 1-5 were determined. In the A2780 test, compound 3 demonstrated considerable cytotoxic action, with an IC50 of 0.7 microg/ml [11]. Through a bioassay-guided fractionation, the cytotoxic chemical moronic acid (1) and the novel tetracyclic triterpene 3,4-seco-olean-18-ene-3,28-dioic acid (2) were obtained from the aerial portions of the medicinal plant Phoradendron reichenbachianum (mistletoe, Loranthaceae). This plant species also contains squalene, glycerol trilinoleate, morolic acid, and flow cytometry investigations indicate that moronic acid, present in galls of Rhus chinensis and Brazilian propolis, at 10microM suppresses the production of Rta, Zta, and an EBV early protein, EA-D, after lytic induction with sodium butyrate. This study also reveals that moronic acids decreases the capacity of Rta to activate a promoter that contains a Rta-response element, showing that moronic acid interferes with the function of Rta. On the other hand, moronic acid does not appear to impact with the transactivation function of Zta. Therefore, the absence of expression of Zta and EA-D following moronic acid treatment is related to the suppression of the transactivation activities of Rta. Because the expression of Zta, EA-D and many EBV lytic genes rely on Rta, the treatment of P9HR1 cells with moronic acid greatly decreases the quantities of EBV particles generated by the cells following lytic induction. This study shows that moronic acid is a novel structural lead for anti-EBV medication development [8].

Anti-HIV

Rhuscholide A (1), a novel benzofuran lactone, was isolated from the stems of Rhus CHINENSIS, along with six other compounds: 5-hydroxy-7-(3,7,11,15-tetramethylhexadeca-2,6,10,11-tetraenyl)-2(3H)-benzofuranone (2), betulin (3), betulinic acid (4), morionic acid (5), 3-oxo-6-beta-hydroxyolean-12-en-28-oic acid (6), and 3-oxo-6-beta-hydroxyolean-18-en-28-oic acid (6) [7]. The structure of rhuscholide A was determined using 1D, 2D NMR (COSY, HMOC, HMBCC) and mass (EI-MS, HR-MS) spectrum data (propan-2-ylidene)-7-(3,7,11,15-tetramethylhexadeca-2,6,10,11-tetraenyl)-2(3H)-benzofuranone [1]. Compound 1 has strong anti-HIV-1 activity, with an EC50 value of 1.62 microM and a therapeutic index (TI) of 42.40, according to in vitro anti-HIV-1 bioassays. With EC50 values of 3.70, 5.81, 7.49, and 11.31 microM, compounds 2, 4, 5, and 7 demonstrated moderate anti-HIV-1 activity [9]. Morolic acid (1) is a pentacyclic triterpene found in nature, and its derivatives have anti-HIV and other biological properties. Starting with betulin, an efficient synthesis of 1 was completed in 11 stages with a total yield of 24%. Moradilol (4), acridocarpic acid D (5), acridocarpic acid E (6), and moronic aldehyde (7) are some of the related natural triterpenes that have been produced. The biological assays revealed that 1, 5, and 6 had a modest inhibitory effect on glycogen phosphorylase [10].

Cytotoxic activity

Five new triterpenoids, acridocarpic acids A-E (1-5), three known triterpenoids, moronic acid (6), ursoic acid, and oleanolic acid, and two known flavonoids, 4',5-dihydroxy-7-methylflavone and 4',5-dihydroxy-3',7-dimethylflavone, were isolated from the cytotoxic MeOH extract obtained from Acridocarpus. On the basis of thorough 1D and 2D NMR spectroscopic data interpretation, the structures of the novel compounds 1-5 were determined. In the A2780 test, compound 3 demonstrated considerable cytotoxic action, with an IC50 of 0.7 microg/ml [11]. Through a bioassay-guided fractionation, the cytotoxic chemical moronic acid (1) and the novel tetracyclic triterpene 3,4-seco-olean-18-ene-3,28-dioic acid (2) were obtained from the aerial portions of the medicinal plant Phoradendron reichenbachianum (mistletoe, Loranthaceae). This plant species also contains squalene, glycerol trilinoleate, morolic acid, betulinaldehyde, betulinaldehide, alpha-germanicol, lupenol, beta-sitosterol, and beta-sitosterol glycopyranoside. Chemical and spectroscopic data were used to deduce the structures [12].

Anti-herpes

Rhus javanica, a medicinal plant, has been demonstrated to have anti-HSV action in mice. We isolated two main anti-HSV chemicals, moronic acid and betulinic acid, from the herbal extract and tested their anti-HSV efficacy in vitro and in vivo. It was the main anti-HSV component in the ethyl acetate fraction. The effective concentrations...
of moronic acid and betulonic acid for 50% plaque reduction were 3.9 and 2.6 microgram/ml, respectively. Betulonic acid had a higher therapeutic index (10.3-16.3). (6.2). HSV-1 resistant to acyclovir-phosphonoacetic acid, thymidine kinase-deficient HSV-1, and wild-type HSV type 2 were all susceptible to moronic acid. When given orally to mice infected cutaneously with HSV-1 three times daily, this chemical substantially delayed the development of skin lesions and/or lengthened mean life periods without harm. Moronic acid suppresses viral production more effectively in the brain than the skin. This was in line with longer mean survival periods. Moronic acid was isolated as a significant anti-HSV component from Rhus javanica. ACV's anti-HSV action differed from ACV's. Moronic acid and betulonic acid for 50% plaque reduction were against Gram-positive bacteria. The active component C30H46O3 was Ozoroa mucronata root bark extract demonstrated antibacterial action used herb, there are no research to support it [15].

Antimicrobial activity

Ozoroa mucronata root bark extract demonstrated antibacterial action against Gram-positive bacteria. The active component C30H46O3 was isolated from the extract during bioassay. Spectroscopic investigations revealed the first natural olean-18-ene keto acid structure 1 isolated from the extract during bioassay. Antimicrobial activity infected mice [13].

Poria cocos fungus and moronic acid from propolis is on the therapeutic index (10.3 -16.3). (6.2). HSV -1 resistant to acyclovir -of moronic acid and betulonic acid for 50% plaque reduction were against Gram–positive bacteria. The active component C30H46O3 was Ozoroa mucronata root bark extract demonstrated antibacterial action

CONCLUSION

Moronic acid is a pentacyclic triterpene derived from the sumac plant Rhus javanica, which has long been thought to have therapeutic properties. Various studies on its pharmacological effects have sparked interest in health-promoting characteristics like as Antidiabetic activity, Anti-AIDS agents, Chemotherapeutic agents, Virus lytic, Anti-HIV, Cytotoxic activity, Anti-herpes, Antimicrobial activity, Ribosome-loaded mRNAs.

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Authors Contributions

All the authors have contributed equally.

Conflits of interests

Declared none

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