Antiviral Properties of Baicalin: a Concise Review

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Received: 8 July 2021 / Accepted: 26 August 2021 / Published online: 6 October 2021
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Abstract
Baicalin is one of the bioactive flavonoid glycosides isolated from the dried root of Scutellaria baicalensis Georgi, Lamiaceae, with antiviral properties. In recent years, the antiviral activity of baicalin has been widely investigated to explore its molecular mechanism of action. In this mini-review, the molecular mechanisms of action of baicalin as an antiviral agent are evaluated, which included three categories: the inhibition or stimulation of JAK/STAT, TLRs, and NF-κB pathways; up or down modulation of the expression levels of IFN, IL, SOCS1/3, PKR protein, Mx1 protein, and AP-1 protein; and inhibition of cell apoptosis caused by virus infection. In addition, clinical studies of baicalin are also discussed. This literature search suggested that baicalin can serve as a potential candidate for the development of a novel broad-spectrum antiviral drug.

Keywords Flavonoid glycoside · Bioavailability · Molecular mechanism · Signaling pathway · Toxicity · Clinical studies

Introduction
In recent years, the public health security has attracted wide attention from people all over the world, particularly after COVID-19 pandemic around the world. Given the scarcity of efficacious and safe vaccines or drugs for coronavirus disease 2019, natural products are a non-orthodox therapeutic approach in the management of the disease that has gained attention, especially in developing countries (Chinsembu 2020; Komolafe et al. 2021). Several viruses cause a biochemical imbalance in the infected cell resulting in oxidative stress that promotes cellular damage through protein denaturation, changes in the functions of nucleic acids, lipid peroxidation, and cell death. These effects may be associated with the development of pathologies and worsening of symptoms. Antioxidant properties of polyphenolic compounds (active redox agents), such as flavonoids, have been associated with the presence of aromatic phenolic rings that promote the electron donation and hydrogen atom transfer to free radicals, acting as free radical scavengers, reducing agents, and quenchers of single oxygen formation that can be used as antiviral agents (Diniz et al. 2020).

Baicalin (1) is the main flavonoid glucoside of Scutellaria baicalensis Georgi, Lamiaceae, which has antiviral, bacteriostatic, antitumor, and anti-oxidative effects (Long et al. 2019). This medicinal plant, known with the common name of Baikal skullcap or Chinese skullcap, is native to China, Korea, Mongolia, and in the Russian Far East and Siberia. It is one of the 50 fundamental herbs used in the traditional Chinese medicine (TCM), where it has the name of huáng qín (Chinese: 黃芩). This plant was originally documented in Shenmeng Bencao Jing, the first Chinese Materia Medica during the Eastern Han Dynasty (25 CE–220 CE) and described the crude dug as the dried root with a bitter taste and cold in nature (Sun and Sun, 1995). Medicinal values of the Chinese skullcap include heat-clearing, elimination of dampness, detoxification, and bleeding prevention (Fang et al. 2020). This plant is used for the prophylaxis and treatment of hepatitis, atherosclerosis, dysentery, ulcerative colitis, inflammations, metabolic disorders, hypertension, hyperlipidemia, type 2 diabetes, and bacterial and viral infections in the respiratory and gastrointestinal tracts (Dinda et al. 2017; Liang et al. 2019; Fang et al. 2020; Su et al. 2020). Baicalin (1) is regarded as the therapeutic chemical marker for quality control of the S. baicalensis roots (Tong et al. 2012). To improve the therapeutic effects,
different herbs are usually combined to produce the TCM formula. The interactions of herbs in the prescription may alter the pharmacokinetics of baicalin. As a major component of TCM formulas, *S. baicalensis* is widely prescribed with herbs such as *Coptidis rhizoma* (Lou et al. 2021) and *Bupleuri radix* (Huang et al. 2019).

Baicalin (I) has been found to have antiviral effect on SARS-CoV-2 (2019-nCoV) *in vitro* (Chen et al. 2020; Jo et al. 2020; Russo et al. 2020; Su et al. 2020). Baicalin (I) and baicalein, its aglycone, were characterized as the first noncovalent, non-peptidomimetic inhibitors of SARS-CoV-2 3CL protease, which is indispensable for coronavirus replication, and exhibited potent antiviral activities in a cell-based system (Su et al. 2020). With increasing interest in the antiviral effect of baicalin, many studies have been carried out focusing on its pharmacological actions. We have summarized baicalin’s mechanism of action reported in the literature in different cytokines and cell pathways. The pharmacological value of baicalin (I) for the development of a broad-spectrum antiviral drug is further discussed.

**Search Strategy**

A systematic search of the literature was performed in PubMed, Web of Science, China National Knowledge Infrastructure, Scopus, Embase, Google Scholar, and SciFinder Scholar database. Search on baicalin was done by using combinations of keywords, including “Huang Qin,” “baicalin,” “Scutellaria baicalensis Georgi, Labiatae,” “pharmacological activity,” “Physicochemical Properties,” “Pharmacokinetics,” “toxicity,” “quality control,” and “clinical study.” Considering the language limitation, this review only refers to Chinese and English texts. In the present study, the existing animal and clinical evidence regarding the therapeutic effects of baicalin on antiviral will be reviewed in addition to its possible mechanism.

**Discussion**

**Physicochemical Properties**

Baicalin (I) is practically insoluble in water (0.054 g/l) and soluble in alcohol, with a molecular weight and melting point of 446.4 g/mol and 222–224 °C, respectively. The basic structure of baicalin corresponds to a flavonoid, a benzopyran ring with a phenolic moiety, as the aglycone with a glucuronic acid moiety forming an anomeric linkage with the C-7 hydroxyl position of the aglycone which contributes to the solubility of the molecules in polar protic solvents. Log P 1.1 indicates that the molecule is poorly lipophilic. The presence of a di-hydroxylated (catechol) substitution on ring A is the main property of the active redox polyphenolic compounds. Catechols in flavonoids and polyphenols are able to form chelates with metals and are reactive in their oxidized form (like quinonas) for radical scavenging and enzyme inhibition with nucleophiles present in lateral protein chains such as cysteine and lysine.

Baicalin (I) is unstable in the basic pH and stable in weakly acidic pH. Baicalin contains two phenolic hydroxyl and one carboxyl groups. Therefore, the acid property of baicalin was characterized in terms of three acid dissociation constants (Kₐ): 5.05, 7.6, and 10.1 (Liang et al. 2009). It is reported that baicalin has poor water solubility (67.03 ± 1.60 μg/ml), low permeability (Papp = 0.037 × 10⁻⁶ cm/s) and low absolute bioavailability (2.2 ± 0.2%) (Xing et al. 2005; Wu et al. 2014). Thus, one of the challenges with baicalin is to improve its bioavailability.

**Quantification**

Several analytical methods, mainly HPLC and UPLC coupled to UV, ECD, MS, and MS/MS, were reported for the quantitative determination of baicalin (I) and its related glycosides in biological samples. Li et al. (2011) reported the HPLC–UV method to quantify the content of baicalin, their metabolites in rat’s blood serum after oral administration of the products containing *S. baicalensis*. It is also reported that the absorption of baicalin occurs at two sites: in upper intestine in the form of glycoside and in the colon in the form of aglycone (Lu et al. 2007). Furthermore, HPLC–ECD (Kotani et al. 2006) and UPLC–MS (Wang et al. 2009) methods were developed for the qualitative and quantitative determination of baicalin, puerarin, wogonoside, and liquiritin in rat’s blood. This method is being used to determine the quality of Gegen-qinlian Tang and other *S. baicalensis* products. By application of a LC–MS method, it was reported that *xiao-chai-hu tang*, a herbal formula for chronic hepatitis B containing *S. baicalensis* (Kong et al. 2018), has a significant influence on the absorption and elimination of baicalin and wogonoside (Zhu et al. 2010). A LC–MS/MS method was reported for determining the analysis of the pharmacokinetics of compounds in rats upon the oral administration of various *S. baicalensis* preparations (as per Chinese Materia Medica) (Kim et al. 2006; Feng et al. 2010; Ye et al. 2010).
Bioavailability and Pharmacokinetics

Bioavailability

Baicalin (1) is categorized as class IV drug under BCS classification. Due to the glycosidic nature of this flavone, baicalin is too polar to cross the lipid bilayer by passive diffusion and is thus poorly absorbed in the intestinal tract. Conversely, baicalein, the aglycone of 1, demonstrates a good permeability due to its great lipophilicicy and is well absorbed in the gastrointestinal tract. Baicalin is moderately absorbed in the stomach but poorly in the small intestine and colon, while baicalein had better absorption than baicalin in all segments of the gastrointestinal tract. However, baicalin, rather than baicalein, is the major component in the systemic circulation following oral administration of baicalein (Liu and Jiang 2006).

To improve its bioavailability, several formulations such as cyclodextrin complexes, self-nanoemulsifying systems, micelles, nanocrystals, and gels were developed (Jakab et al. 2019; Xing et al. 2005). It was reported that the efflux of baicalin might be mediated by ATP-binding cassette (ABC) transporters (Kalapos-Kovács et al. 2015). Jakab et al. (2019) and Li et al. (2018) have developed cyclodextrin complexes for baicalin via the inclusion complexation by supercritical fluid encapsulation technique to improve the bioavailability of this flavonoid glucoside. Additionally, owing to hydrogen bonding and van der Waals interactions, the coordination complex of baicalin with the oxovanadium (IV) cation, Na4[VO(baic)2]·6H2O VO4(baic), has stronger binding with bovine serum albumin than baicalin (Martínez Medina et al. 2017). A self-emulsifying microemulsion drug delivery system (SMEDDS) was developed for baicalin using a phospholipid complex (PC) by a solvent evaporation method. The formulation has improved the bioavailability, and the relative bioavailability of baicalin–PC (1:2)–SMEDDS was 220.37% of baicalin in rats (Wu et al. 2014). Furthermore, the nanoemulsion was reported to improve the absorption of baicalin from the intestinal tract compared to baicalin suspension. The absorption rate constant (Kα) and apparent permeability coefficient (Papp) were increased by 6.14-fold and 6.25-fold, respectively (Wu et al. 2018). In addition, baicalin nanoemulsions were reported to increase the saturation solubility of baicalin and strengthened their interaction with the mucin layer in epithelial tissues leading to an increased uptake in the intestine. The AUC_{(0-1)} and C_{max} of baicalin from a lipid-based nanoemulsion was reported to be significantly higher (11.5-fold) than baicalin powder for lymphatic system absorption (Xu et al. 2019). The in vivo disposition of baicalin is affected by combinations of other herbs, and baicalin can interact with other co-administered drugs due to competition between metabolic enzymes and protein binding (Huang et al. 2019). Some other studies have reported the development of nanocapsules and nanocrystals which have improved the absorption and bioavailability of baicalin in pre-clinical studies (El-Gogary et al. 2019; Jin et al. 2014).

Pharmacokinetics

Baicalin (1) is a flavone glycoside mainly absorbed in the stomach but poorly absorbed in the small intestine and colon in comparison to its aglycone, baicalein (Liu and Jiang 2006). Baicalin displays a distinct pharmacokinetic profile including gastrointestinal hydrolysis, enterohepatic recycling, carrier-mediated transport, and complicated metabolism. Numerous studies have reported that the enzymes, β-glucuronidase and UDP-glucuronosyltransferase (UGT), play a vital role in the hydrolysis of baicalin to baicalein (Fong et al. 2012; Kalapos-Kovács et al. 2015; Noh et al. 2016). It is interesting to note that baicalin mainly exist in its glycosidic form in systemic circulation (Li et al. 2014). The studies have observed bimodal and multiple peaks in the absorption profile of baicalin (Zhao et al. 2014; Zhang et al. 2016). These peaks probably arise due to variable gastric emptying time, glucuronidation, and absorption at different sites (Lu et al. 2007; Tong et al. 2012; Song et al. 2017). In addition, after combining with human plasma protein, baicalin was reported to bind with serum albumin quickly and was distributed to various tissues mediated through ABC transporter inhibitors in membranes overexpressing the ATP-dependent MDR1 efflux pump in Caco-2 cell monolayer model (Zhang et al. 2007). Additionally, the concentration of baicalin is different in various visceral organs depending on the formulation and route of administration (Zhu et al. 2013; Wei et al. 2016). The metabolites of baicalin have been observed in various tissues (heart, liver, spleen, lung, kidney, and brain), and the main organs for its metabolism were reported to be the liver and kidney (Zhang et al. 2015). The predominant enzymes involved in metabolism of baicalin are UGT, catechol-O-methyltransferases, β-glucuronidase, and sulfatase and thus causes methylation, methylation, hydroxylation, sulfation, glucuronidation, hydrolysis, etc. (Akao et al. 2013; Lu et al. 2012; Wang et al. 2012; Zhang et al. 2015). Baicalin was reported to be predominantly excreted in bile which is mediated by MRP2 (Xing et al. 2005; Akao et al. 2009; Kalapos-Kovács et al. 2015). It was reported that only 7.2% of the baicalin was primarily excreted in urine in the form of sulfated and hydroxylated derivatives and parent drug (Jiang et al. 2014; Lai et al. 2003).

Toxicity

A recent study has reported that baicalin (1) and baicalein inhibit the main protease (3CLpro) of SARS-CoV-2 (Su et al. 2020). Very few studies have been conducted...
cytes and granulocytes were observed (Han et al. 2011). Lung alveolar fusion, and increased number of monocytic cells (RBL-2H3) without causing dermal irritation/corrosion (Kim et al. 2013). In a 90-day oral toxicity study of UP446 (a combination of baicalin from the roots of Scutellaria baicalensis and (+)-catechin from the heartwoods of Acacia catechu), the ‘No Observed Adverse Effect Level (NOAEL)’ was found to be 1000 mg/kg/day (Yimam et al. 2010). Another study has confirmed that a single-oral dose of 100–2800 mg of baicalin is safe and well tolerated in healthy volunteers (clinical trial registration number: CTR20132944) (Li et al. 2014). However, a recent study has demonstrated that baicalin can activate the transforming growth factor-β (TGF-β)/Smad signaling pathway in a dose-dependent manner, induce the Smad3 acetylation (via interaction of Smad3 with transcriptional coactivator p300), and reduce the phosphorylation of AMPK (a metabolic master kinase) that results in kidney injury and fibrosis at higher doses (Cai et al. 2017). Han et al. (2011) evaluated in vitro and in vivo the pulmonary toxicity of shuang-huang-lian preparation, with a long history for treating respiratory tract infection. A report indicated that the injection of the lyophilized powder of shuang-huang-lian prepared with honeysuckle (Lonicera caprifolium L., Caprifoliaceae), Scutellaria baicalensis roots, and the fruits of Forsythia suspensa (Thunb.) Vahl, Oleaceae (Dong et al. 2017), is commonly used to treat bacterial and viral infections in the respiratory tract (Zhang et al. 2013). The in vitro results showed no adverse effects at doses of up to 6 mg/kg and 25 mg/kg for histopathologic study and cell counts experiment, respectively. In vivo results showed no difference in LDH activity, total protein content, nor IL-6, IL-8, and TNF-α levels in bronchoalveolar lavage fluid (BALF). However, inhalation of shuang-huang-lian preparation at higher doses causes lung alveolar fusion, and increased number of monocytes and granulocytes were observed (Han et al. 2011).

### Antiviral Activity

#### Signaling Pathway

**IFN and JAK-STAT** Interferons (IFNs) are a group of signaling proteins of cytokine which are secreted by the human body after virus infection and are named for their ability to “interfere” with viral replication by the regulation of the immune system (Darnell et al. 1994). In pulmonary microvascular endothelial cells (PMVEC), the secretion of IFN-α and IFN-γ is significantly increased to reduce the degree of immunological injury (Lin et al. 2009). Therefore, the effect of baicalin (I) on IFN expression has been explored as an important molecular mechanism for the antiviral effect of this flavonoid. The JAK-STAT signaling pathway is involved in processes such as immunity, cell division, cell death, and tumor formation. There are three key parts of JAK-STAT signaling: Janus kinases (JAKs), signal transducer and activator of transcription proteins (STATs), and receptors (which bind the chemical signals). It has been reported that baicalin exhibited antiviral effect as demonstrated by the use of the respiratory syncytial virus infection model via the following mechanism: (1) the increasing expression of IFNs in rats (Zhan et al. 2017), (2) activation of JAK-1 and non-receptor tyrosine-protein kinase (TYK-2) to promote phosphorylation of STAT1 and STAT2, and (3) inhibiting expression of SOCS1 and SOCS3 proteins to activate JAK/STAT pathway (Zhang et al. 2018).

Numerous studies claimed that baicalin (I) could help to induce IFN-γ secreted by CD4+, CD8+ t-lymphocytes, and NK cell to effectively enhance activity of JAK/STAT1 pathway (Chu et al. 2015; Zhi et al. 2019). The main chemical components of the shuanghuanglian injection are chlorogenic acid, baicalin, and forsythia glycosides, which have been found to have the ability of anti-inflammation, improving immunity, and inhibiting the growth of various viruses. Baicalin, as one of the bioactive ingredients, is capable of increasing expression of IFN-γ and effectively inhibiting cell apoptosis in a dose-dependent and time-dependent manner with infection of human adenovirus III (HAdV3) in A549 and Hep-G2 cells (Ma et al. 2017). Oral administration of Scutellaria baicalensis root (80% ethanol) effectively protected mice infected with influenza A virus (IVA), increased the survival rate (herbal preparation: 67%; baicalin: 33%), decreased the lung injury index (herbal preparation: 0.90; 1: 1.00), and improved the lung morphology (Fig. 1). This preparation efficiently decreased lung virus titers, reduced hemagglutinin titers, inhibited neuraminidase activities in the lungs of IAV-infected mice, and modulated the inflammatory responses by

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decreasing the levels of tumor necrosis factor-α (TNF-α), interleukin-6 (IL-6), and monocyte chemotactic protein-1 (MCP-1) and increasing the levels of interferon-γ (IFN-γ) and interleukin-10 (IL-10) in lung tissues (Zhi et al. 2019).

**TLRs and NF-kB** NF-κB (nuclear factor kappa-light-chain-enhancer of activated B cells) is a protein complex that controls transcription of DNA, cytokine production, and cell survival. NF-κB is found in almost all animal cell types and is involved in cellular responses to stimuli such as stress, cytokines, free radicals, heavy metals, ultraviolet irradiation, oxidized LDL, and bacterial or viral antigens (Fig. 2). NF-κB induced immunological injury through regulating the expression of TAB2, NFKB, IRAK1 and RIPK1 (Liu et al. 2016). A report suggested that baicalin (1) inhibit immunological injury through the regulation of TNF-α, IL-1β, IL-6, IL-8, IL-12, IL-18, and CCL5 via specifically inhibiting TLR7/MyD88 pathway. In this pathway, the expression of TLR3, TLR4, and TRIF mRNA were suppressed; the levels of IRAK4 and FOS were reduced; NF-κB and AP-1 protein were activated; and the phosphorylation of NF-kB-65 protein was decreased (Wan et al. 2014). These observed changes suggest that the primary mechanism of baicalin (1) in antiviral in inhibiting immunological injury involves inhibiting NF-kB and TLR7/MyD88 pathway, followed by reducing the inflammatory cytokine expression.

**HNFs** HNFs plays a vital role in the proliferation of hepatitis B virus (HBV) and the underlying mechanism involved in up-regulating HNF1α expression, increasing HNF4α expression, and promoting pgRNA biosynthesis and transcription (Huang, 2013). In Hep-G2 cells, baicalin (1) substantially decreased HBsAg and HBeAg expression via reducing the level of HNF1α and HNF4α to inhibit the RNA of HBV proliferation and synthesis (Cheng et al. 2006; Huang et al. 2017b). In addition, a study in exploring the mechanisms involved in the treatment of HBV has been reported for baicalin to reduce proliferation through increasing the physical interaction of HNF4α and ERα/β and decreasing the expression of HNF1α to inhibit the bond between HNFs and cccDNA (Chirumbolo 2018; Xia et al. 2020).

**Interferon Regulatory Factors** Influenza A virus (IAV) causes immunologic injury via mediating inflammatory cytokines. The underlying mechanisms probably involved the regulation of interferon factors (IRFs, such as IRF3 and IRF7), NF-kB activation via promoting retinoic acid-induced protein I (RIG-I) and MAVS protein interaction, and increased IFN-α/β and TNF-α secretion. A recent study has reported that baicalin (1) increases the expression of fundamental factors (IRF3, IRF7, and NF-kB) in RLRs pathway. So, it is suggested that baicalin can suppress innate immune hyperactivity caused by IAV infection (Pang et al. 2018). In ANA-1 cell, the level of macrophage is significantly increased after H1N1 infection, which probably is one of the most important mechanisms of promoting inflammatory reaction occurrence. Thus, decreasing the secretion of macrophage is an effective method to treat inflammatory reaction. A recent study has proven that baicalin increased M1 macrophage ratio, stimulates IFN-β and IRF3 expression, and reduces IL-10, thus inhibiting inflammatory reaction (Geng et al. 2020). Moreover, the chick embryo fibroblast (CEF) cell is reported to overexpress TRF7 after Marek’s disease virus (MDV) infection. Yang et al. (2020) reported that baicalin decreased the IRF7 level in CEF cell, thus exhibiting antiviral effects.

**Interleukin (IL) and TNF-α** When the virus invades the host body, a series of physiological reactions occur in the host body, including the activation of the immune response and the secretion of a variety of inflammatory factors. IL-6, a common inflammatory cytokine, plays a vital role in increasing the expression of STAT3 protein and SOCS-1 gene and inhibiting the differentiation of Th1 cell, thus down-regulating the JAK/STAT pathway (Zhang et al. 2018). Zhang et al. (2020) reported that baicalin (1) has shown a significant effect in relieving oxidative stress by reducing the level of IL-17A, IL-22, IL-6, and IL-1p in chicks infected with H9N2. Treatment of baicalin lowered the TNF-α and IL-2 level and reduced the degree of inflammatory cell infiltration in RSV-infected rat (Cheng et al. 2014). Moreover, inflammatory injury is caused by human adenovirus III in A549 and Hep-G2 cell. After being treated by shuanghuanglian injection lyophilized powder, the symptom was significantly reduced. The underlying mechanisms were reported to the inhibition of plaque formation and decreasing the levels of TNF-α, IL-1β, and IL-6 (Ma et al. 2017).

**Apoptosis-Related Genes**

The third phase of cell apoptosis, center control and effective phase, is the phase of cysteine aspirate protease activated by apoptosis signal to performed programmed cell death. The apoptosis pathways include caspase-8-mediated extrinsic pathway and the intrinsic pathway (caspase-9-mediated mitochondria apoptosis pathway and endoplasmic reticum pathway) (Long et al. 2004; Nomura et al. 2005). In recent years, studies reported that baicalin (1) exhibits the inhibition of cell apoptosis via (1) decreased expression of membrane receptor Fas; (2) reduced level of caspas; (3) increased expression of Bcl-xl, FLIP, and IAP; (4) inhibition of protein phosphatase 3 that catalyzes the subunit alpha-subtype gene (Cn); and (5) inhibition of apoptosis signal transduction. Therefore, the result suggested that baicalin exhibited inhibition of cell apoptosis through regulating the relevant protein of caspase-8.
expression mediated by extrinsic pathway and endoplasmic reticulum pathway. In addition, in the cell infected with EV71 and CHIKV, the expression of pro-apoptotic genes, Fas and caspase-3, is significantly decreased to inhibit cellular apoptosis. Therefore, the effect of baicalin (1) on cellular apoptosis has been explored as a vital underlying mechanism for the antiviral effect of this flavonoid (Li et al. 2015b; Oo et al. 2018).

**Immune Cells**

The activity of the T cell subset is an index to determine the cellular immune function of patients. Recent studies have indicated that baicalin (1) can improve patient immune function. In infected patients with HBV and RSV, baicalin increased CD4+, CD8+ levels, CD4+/CD8+ ratio, and the activity of NK cells (Shi et al. 2016). The IAV infection was reported to increase elevated level of pro-inflammatory response factors (Th1 and Th17). Baicalin was reported to inhibit the inflammatory reaction via increased up-regulation of the Th2 and Treg expression and decreased Th1/Th2 and Th17/Treg ratios. The results revealed that baicalin has blocked the Th1 and Th17 type inflammatory reaction through regulating the differentiation of CD4+ (Pang et al. 2018). Baicalin was reported to inhibit H1N1 infection via stimulation of CD4+, CD8+, and NK cells, increased

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**Fig. 1** Hematoxylin–eosin stained (400×) of lung tissues (A) and immunohistochemical expression of C3d in lungs (magnification of alveolus) at day 4 after infection (B). Abbreviations: Model, influenza A virus-infected lung tissues; Ext, extract of *Scutellaria baicalensis* root in 80% ethanol; 1, baicalin; RB, ribavirin (positive control). Reproduced from Zhi et al. 2019
expression of IFN-γ, and activation of JAK/STAT1 pathway in human peripheral blood monocyte cells (HPBMC) (Chu et al. 2015).

Others

In Hela cell, Coxsackie Virus B3-induced cell autophagy is mediated via (1) increasing the content of free fatty acid (FFA) and cholesterol (CHO), (2) up-regulating the expression of fatty acid synthase (Fasn) and acetyl-CoA carboxylase mRNA, and (3) stimulating the generation of cytolsosomes. The treatment of baicalin (1) reduced the level of FFA, CHO, Fasn acetyl-CoA carboxylase, and LC3-II (an autophagosome marker). Therefore, the result has concluded that the observed antiviral effect of baicalin on CVB3 might be mediated via suppressing lipid synthesis and inhibiting autophagosome generation (Wang et al. 2020). The mechanisms implicated in the inhibitory effects of baicalin on viral pneumonia were also reported in the literature. In FM1-infected mouse model, baicalin was reported to down-regulate the MDA and total NO synthase levels in the lung tissues, increase the glutathione peroxidase (GSH-PX) activity, reduce endothelial cell dysfunction, reduce the vascular overexpansion, and reduce the tissue injury (Wan et al. 2011). Finally, the oral liquid shuang-huang-lian formulation showed a dose-dependently inhibited SARS-CoV-2...
3CLpro as well as the replication of SARS-CoV-2 in Vero E6 cells. Baicalin and baicalein were characterized as the first noncovalent, nonpeptidomimetic inhibitors of SARS-CoV-2 3CLpro and exhibited potent antiviral activities in a cell-based system (Su et al. 2020).

To sum up, the most significant antiviral molecular mechanisms of baicalin (1) are the increased expression of IFNs; activation of JAK/STAT pathway; suppression of TLRs activity; suppression of NF-κB pathway; reduction of HNFs, IRFs, TNF-α, and IL levels; inhibition of cysteine aspirate protease; inhibition of pro-apoptotic genes; and improvement of immune function. The antiviral molecular mechanisms of baicalin are summarized in Table S1, and the molecular pathways for the antiviral properties of baicalin are illustrated in Fig. 3.

Clinical Studies

With the rising interest in the antiviral effect of baicalin (1), numerous studies have been carried out focusing on its clinical applications. Baicalin could be combined with other medicines to treat viral diseases such as chronic hepatitis B and A H1N1 influenza. The possible underlying molecular mechanisms involved in its beneficial action are improving the immunity in patients via regulation of the T-lymphocyte subset levels and up-regulation of CD3+, CD4+, and CD4+/CD8+ levels in patients infected with A H1N1 influenza (Su and Xiang 2014). In addition, a study has reported that baicalin reduces the level of alanine aminotransferase (ALT), aspartate transaminase (AST), and total bilirubin (TBil), and HBV-DNA were reduced and reduced the hepatic fibrosis in chronic hepatitis B patients (Huang et al. 2017a). The details about drugs and studies are summarized in Table S2.

However, the reported findings in the literature are obtained from retrospective studies, and thus there might be a component of bias in reporting the efficacy. The bias could arise from the subjectivity of the clinician, experimental methods, sample size, and the classification of disease, which are parameters difficult to control in clinical studies (Li et al. 2020, 2021).

Perspective on Future Directions

Some clinical trials have suggested that the potent inhibitory activity of baicalin (1) on influenza virus make it worthwhile to be subjected to further clinical trials, particularly for COVID-19 treatment (Su et al. 2020). Low aqueous solubility and low bioavailability of baicalin (1) are the major challenges for its use in clinic. Encouraging results in terms of improving the bioavailability are obtained from cyclodextrin complexes, self-nanoemulsifying systems,
micelle, nanocrystals, and gelling system, *inter alia*. However, development of an ideal formulation is still far away. In the literature, very limited studies were reported on the safety and efficacy of baicalin. Also, a few clinical studies have reported the beneficial effect of baicalin in improving the antiviral activity of entecavir, lamivudine, and oseltamivir, which has to be further evaluated for the treatment of viral infections in the respiratory tract and combat COVID-19. There were only very few clinical studies in which the antiviral activity of baicalin was investigated. Therefore, the future research studies must be focused on evaluating the clinical efficacy of baicalin in treating viral infections. Therefore, more rigorous scientific studies should be carried out on baicalin and its plant sources.

**Conclusion**

In this review, we have discussed the latest research progress on baicalin (I) regarding its physicochemical properties, toxicity, bioavailability, pharmacokinetics, antiviral activities, and clinical studies. In recent years, the number of studies on baicalin was increased, and most of the studies have been concentrated on the direction of the clinical studies for its effect as an antiviral activity. Additionally, despite the availability of many reported formulations on baicalin to improve its bioavailability, the ideal formulation is yet to be identified. Only a few clinical trials with small number of volunteers have been conducted to evaluate the safety of baicalin, and thus utmost care for the administration of baicalin should be paid. In this review, we also summarized the antiviral molecular mechanisms of baicalin. In recent years, the antiviral effect of baicalin against IVA, RSV, HV, CVB3, etc. has been widely investigated. The main molecular mechanisms reported were regulation of JAK/STAT, TLRs, and NF-κB pathways. Baicalin inhibits inflammations via regulating the IFN, IL, SOCS1/3, PKR protein, Mx1 protein, and AP-1 protein levels. The antiviral activity of baicalin was also reported to be exerted through inhibition of apoptosis, decreased expression of FAS protein, inhibition of caspase-8 pathway, and decreased expression of Bax protein. These information present in this review will be helpful in guiding future research studies. Further studies must be carried out to explore the possibility of involvement of additional mechanisms.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s43450-021-00182-1.

**Acknowledgements** The authors thank the Shandong University of Traditional Chinese Medicine Library for providing access to ancient Materia Medica books.

**Author Contribution** KL and YL conceived and designed the review. AC, QW, YL, and HW collected and analyzed literature information. KL and YL wrote the manuscript. CZ and XW reviewed the manuscript. All authors have read the final manuscript and approved its submission.

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