Antiplatelet and antithrombotic effects of cordycepin-enriched WIB-801CE from *Cordyceps militaris* ex vivo, in vivo, and in vitro

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**Abstract**

**Background:** A species of the fungal genus *Cordyceps* has been used as a complementary and alternative medicine of traditional Chinese medicine, and its major component cordycepin and cordycepin-enriched WIB-801CE are known to have antiplatelet effects in vitro. However, it is unknown whether they have endogenous antiplatelet and antithrombotic effects. In this study, to resolve these doubts, we prepared cordycepin-enriched WIB-801CE, an ethanol extract from *Cordyceps militaris*-hypha, then evaluated its ex vivo, in vivo, and in vitro antiplatelet and antithrombotic effects.

**Methods:** Ex vivo effects of WIB-801CE on collagen- and ADP-induced platelet aggregation, serotonin release, thromboxane A2 (TXA2) production and its associated activities of enzymes [cyclooxygenase-1 (COX-1), TXA2 synthase (TXAS)], arachidonic acid (AA) release and its associated phosphorylation of phospholipase Cβ3, phospholipase Cγ2 or cytosolic phospholipase A2, mitogen-activated protein kinase (MAPK) [p38 MAPK, extracellular signal-regulated kinase (ERK)], and blood coagulation time in rats were investigated. In vivo effects of WIB-801CE on collagen plus epinephrine-induced acute pulmonary thromboembolism, and tail bleeding time in mice were also inquired. In vitro effects of WIB-801CE on cytotoxicity, and fibrin clot retraction in human platelets, and nitric oxide (NO) production in RAW264.7 cells or free radical scavenging activity were studied.

**Results:** Cordycepin-enriched WIB-801CE inhibited ex vivo platelet aggregation, TXA2 production, AA release, TXAS activity, serotonin release, and p38 MAPK and ERK2 phosphorylation in collagen- and ADP-activated rat platelets without affecting blood coagulation. Furthermore, WIB-801CE manifested in vivo inhibitory effect on collagen plus epinephrine-induced pulmonary thromboembolism mice model. WIB-801CE inhibited in vitro NO production and fibrin clot retraction, but elevated free radical scavenging activity without affecting cytotoxicity against human platelets.

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Background
A species of the fungal genus Cordyceps is known to prescribe for inflammatory and cancer disease [1, 2]. It is reported that cordycepin (3’-deoxyadenosine, Fig. 1a), a major component of Cordyceps militaris, has in vitro antithrombotic effects by attenuating [Ca^{2+}]_i level and thromboxane A_2 (TXA_2) production in collagen-induced human platelet aggregation [3]. However, there is no evidence or report concerning ex vivo and in vivo inhibitory effect of cordycepin or cordycepin-enriched substance on platelet activation.

In this study, to resolve this doubtful point, we prepared cordycepin-enriched WIB-801CE (Compound from 2008 First Project of Bioteam, Whanin Pharm. Co., Ltd., Suwon, Korea), an ethanol extract from Cordyceps militaris-hypha. Next, to observe whether WIB-801CE has endogenous inhibitory effects on platelet activation associated with thrombus formation, we orally administered WIB-801CE to rat, and subsequently investigated the effects on major molecules associated with Ca^{2+} increase [4–7], arachidonic acid (AA) release [4, 6, 8–10], TXA_2 production [4, 5, 8, 11–13] and serotonin release [13–16].

Methods
Materials
WIB-801CE was provided from Whanin Pharmaceutical Corporation (Suwon, Korea). Collagen, adenosine diphosphate (ADP) and thrombin were obtained from Chrono-Log Corporation (Havertown, PA, USA). Serotonin enzyme-linked immunosorbent assay (ELISA) kit was purchased from Labor Diagnostika Nord GmbH & Corporation (Nordhorn, Germany). Pure cordycepin, aspirin, protease inhibitor cocktail, 2,2-diphenyl-1-picrylhydrazyl (DPPH), ascorbic acid (AC), and other reagents were obtained from Sigma Chemical Corporation (St. Louis, MO, USA). Thromboxane B_2 (TXB_2) enzyme immunoassay (EIA) kit, cyclooxygenase-1 (COX-1) fluorescent activity assay kit, lactate dehydrogenase (LDH) cytotoxicity assay kit, ogarel and prostaglandin F_2 alpha (PGF_2a) for TXA_2 synthase (TXAS) assay were purchased from Cayman Chemical (Ann Arbor, MI, USA). Arachidonic acid (AA) release ELISA kit was purchased from Cusabio Biotech Corporation (Wuhan, Hubei, China). Anti-phosphor-cytosolic phospholipase A_2 (cPLA_2) (Ser^{505}), anti-phosphor-phospholipase C_{β3} (PLC_{β3}) (Ser^{537}), anti-phosphor-phospholipase C_{β3} (PLC_{β3}) (Ser^{1105}), anti-phosphor-phospholipase C_{γ2} (PLC_{γ2}) (Tyr^{1217}), anti-phosphor-p^{38} MAPK , anti-phosphor-extracellular signal-regulated kinase (ERK) (1/2), anti-p^{38} MAPK , anti-ERK (1/2) and anti-rabbit immunoglobulin G (IgG)-horseradish peroxidase conjugate (HRP), and lysis buffer were obtained from Cell Signaling (Beverly, MA, USA). Polyvinylidene difluoride (PVDF) membrane was from General Electric Healthcare (Piscataway, NJ, USA). Enhanced chemiluminescence solution (ECL) was from General Electric Healthcare (Chalfont St. Giles, Buckinghamshire, UK). Prothrombin time (PT) assay reagent and activated partial thromboplastin time (APTT) assay reagent were obtained from Fisher Diagnostics (Middletown, VA, USA).

Preparation of WIB-801CE
Cordyceps militaris was cultivated, and culture-solution of Cordyceps militaris-hypha was concentrated up to 50° Brix with a rotary vacuum evaporator (Eyela N3000, Rikakikai Co., Ltd., Tokyo, Japan) at 60 °C. The Brix was measured with refractometer (Atago Co., Ltd., Tokyo, Japan). The concentrate was extracted by extraction-shaker (Cosmos 660, Kyungsco Co., Ltd., Seoul, Korea) for 4 h at 40 °C one time with distilled water/95% ethanol (1:3.5, v/v), which was filtered one time using a filter paper (Advantec No.2). The filtrate was completely concentrated at 40 °C one time with distilled water/95% ethanol (1:3.5, v/v), which was filtered one time using a filter paper (Advantec No.2). The filtrate was lyophilized and stored at -20 °C until used. This was named as cordycepin-enriched WIB-801CE.

Analysis of cordycepin in WIB-801CE with HPLC
WIB-801CE was dissolved with 75% methanol, then analyzed by high performance liquid chromatography (HPLC). An Alliance 2695 liquid chromatography system (Waters Co., Milford, MA, USA), equipped with vacuum degasser, quaternary gradient pump, autosampler and photodiode array detector, was connected to Empower software. A hydrosphere C_{18} column (250 mm x 4.6 mm id, 5 μm, YMC Co., Ltd., Kyoto, Japan) was used at a column temperature of 30 °C. The applied-mobile phase gradient program was 0.01 M KH_2PO_4/methanol (95:5, v/v) at
0 min and held for 5 min; 0.01 M KH₂PO₄/methanol v/v) at 20 min and held for 6 min; 0.01 M KH₂PO₄/methanol (95:5, v/v) at 27 min and held 6 min for chromatographic balance. In this step, 99.8% of methanol was used. The flow rate was at 1.0 mL/min and sample injection volume was 10 μL. The ultra violet detection was operated at 254 nm.

**Animals and administration**

We investigated the ex vivo and in vivo effects of WIB-801CE using rats (Sprague-Dawley, male, 200 g) and Institute of Cancer Research (ICR) mice (male, 18 g, Daehan Biolink Co., Ltd., Chungbook, Korea). Rats for ex vivo experiment and mice for in vivo observation were divided into as follows, respectively: WIB-801CE-nontreated group (control), WIB-801CE-treated group, aspirin-treated group as positive control of in vivo, and warfarin-treated group as positive control of ex vivo.

Animals were acclimatized for a week at a temperature of 24 ± 1 °C and humidity of 55 ± 5%. Before oral administration of substances, all animals were fasted for 12 h, then were fed with standard pellets diet (Purina Inc., Korea) had free access to water. WIB-801CE [200, 400 mg/kg-body weight (BW)] and warfarin (1 mg/kg-BW) for ex vivo experiment were orally administered to the rats one per day for seven days, and WIB-801CE (200, 400 mg/kg-BW) and aspirin (100 mg/kg-BW) for in vivo observation were orally administered to the mice once a day for five days. 200 mg/kg-BW of WIB-801CE is corresponded to the minimum dose that inhibits rat platelet aggregation (data not shown). WIB-801CE, warfarin, and aspirin were
dissolved with distilled water. The experiments were proved by the Ethics Committee for Animal Experiments of Whanin Pharmaceutical Corporation (Suwon, Korea/15-NE-016 for rats, 15-NE-008 for mice).

**Preparation of rat platelet-rich plasma and platelet-poor plasma for ex vivo assay**

After the final respective administration, all rats were fasted for 24 h, then after 2 h of WIB-801CE- and warfarin-administration were anesthetized with 20% urethane before sacrifice according to the method of Zhang et al. [17]. The blood was collected from the abdominal aorta. The blood was anti-coagulated with acid-citrate-dextrose solution (0.8% citric acid, 2.2% sodium citrate, 2.45% glucose), and was centrifuged at 250 × g for 10 min in order to obtain platelet-rich plasma (PRP). In order to remove residual red blood cells and white cells, the PRP was again centrifuged at 125 × g for 10 min. Platelet-poor plasma (PPP) was prepared by centrifuging the part of PRP at 1,300 × g for 10 min.

PRP was used to investigate ex vivo platelet aggregation, TXA2 production, serotonin release, COX-1 and TXAS activities, AA release and protein phosphorylation. PPP was used to investigate ex vivo PT and APTT. The number of platelets in PRP was adjusted with PPP to a final concentration of 5 × 10^8/mL. All of the above procedures were carried out at 25 °C to avoid platelet aggregation from any effect of low temperature.

**Preparation of human PRP and washed platelets for in vitro assay**

To investigate in vitro effects of WIB-801CE and cordycepin on fibrin clot retraction, we used human PRP and washed platelets. PRP from normal healthy human volunteers with informed consent was obtained from the Korean Red Cross Blood Center (KRBC, Changwon, Korea), and its experimental use was approved by the KRBC (Safety Supervisor Team-621-2015.02.26) and the Korea National Institute for Bioethics Policy Public Institutional Review Board (Seoul, Korea/PIRB12-072-01) approved these experiments.

Ex vivo measurement of TXB2

To investigate the effect on TXA2 production, the aggregation was terminated by adding ice-cold 5 mM EDTA and
0.2 mM indomethacin to inhibit subsequent conversion of AA to TXA₂. The amounts of TXB₂, a stable metabolite of TXA₂, were determined using a TXB₂ EIA kit according to the procedure described by the manufacturer.

**Ex vivo Western blot for analysis of protein phosphorylation**
Collagen- and ADP-activated rat PRP was centrifuged for 10 min at 1,300 × g under 4 °C to remove PPP and get platelet pellets. The platelets were suspended twice with suspension buffer (138 mM NaCl, 2.7 mM KCl, 12 mM NaHCO₃, 0.36 mM NaH₂PO₄, 0.49 mM MgCl₂, 5.5 mM glucose, 0.25% gelatin, pH 7.4). The suspended platelets (250 μL) were lysed by adding an equal volume (250 μL) of lysis buffer (20 mM Tris-HCl, 150 mM NaCl, 1 mM Na₂EDTA, 1 mM EGTA, 1% triton X-100, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate (serine/threonine phosphatase inhibitor), 1 mM Na₃VO₄ (ATPase, alkaline and acid phosphatase, and protein phosphotyrosine phosphatase inhibitor), 1 μg/mL leupeptin (serine and cysteine protease inhibitor), and 1 mM phenylmethanesulfonyl fluoride (serine protease and acetylcholinesterase inhibitor), pH 7.5).

Platelet lysates were suspended in their equal volume of sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) buffer (62.5 mM Tris-HCl, 10% glycerol, 1% SDS, 1% β-mercaptoethanol, 0.01% bromphenol blue, pH 6.8), then were boiled to completely denature the proteins for 5 min. Aliquots containing 15 μg of protein from each sample tube were subjected to SDS-PAGE (8%, 15 mm gel) according to the method of Laemmli [20].

Proteins in the gel were transferred to PVDF membrane in the presence of transfer buffer (25 mM Tris-HCl, 192 mM glycine, 20% methanol, pH 8.3). PVDF membrane was washed one time for 5 min with Tris-bufferear saline with tween 20 (25 mM Tris-HCl, 140 mM NaCl, 2.7 mM KCl, 0.1% tween 20, pH 7.4), then was blocked with blocking buffer (25 mM Tris-HCl, 140 mM NaCl, 2.7 mM KCl, 0.1% tween 20, 5% skim milk, pH 7.4) for 1 h at room temperature, and subsequently was washed three times for 5 min.

The protein phosphorylation was observed using Western blotting. The dilutions for 1st antibody (anti-phosphor-cPLA₂, anti-phosphor-PLC₁β₁, anti-phosphor-cPLA₂, anti-phosphor-p³⁸ MAPK, anti-phosphor-ERK, anti-p³⁸ MAPK, anti-ERK) and 2nd antibody (anti-rabbit IgG-HRP) were 1:1,000 and 1:10,000, respectively. The membranes were visualized using ECL. Blots were analyzed using the Quantity One, Ver. 4.5 (BioRad, Hercules, CA, USA).

**Ex vivo determination of AA release**
To investigate the effect on AA release, the aggregation was terminated adding ice-cold 5 mM EDTA and 0.2 mM indomethacin to inhibit subsequent conversion of AA to TXA₂, and centrifuged with 200 × g at 4 °C for 10 min. The supernatants were used for the assay of AA release. AA release was measured with a Synergy HT multi-model microplate reader (BioTek Instruments, Winooski, VT, USA) using AA release ELISA kit.

**Preparation of platelet lysates**
We prepared platelet lysates to determine ex vivo COX-1 and TXAS activities. Collagen- and ADP-activated rat PRP was centrifuged for 10 min at 1,300 × g to remove PPP and get platelet pellets. The platelets were then suspended twice with suspension buffer (138 mM NaCl, 2.7 mM KCl, 12 mM NaHCO₃, 0.36 mM NaH₂PO₄, 0.49 mM MgCl₂, 5.5 mM glucose, 0.25% gelatin, pH 7.4). The suspended platelets in the presence of 1% protease inhibitor cocktail were sonicated ten times in sensitivity 100% for 20 s at 4 °C with a sonicator (HD 2070, Bandelin Electronic, Bandelin, Germany) to obtain platelet lysates. Next, the platelet lysates were centrifuged at 12,000 × g for 15 min at 4 °C to remove cell debris. The supernatant was used to measure COX-1 and TXAS activity.

**Ex vivo measurement of COX-1 activity**
Platelet lysates containing 10 μg of protein were used. COX-1 activity was measured with a Synergy HT multi-model microplate reader (BioTek Instruments, Winooski, VT, USA) using COX-1 fluorescent activity assay kit according to the procedure described by manufacturer.

**Ex vivo measurement of TXAS activity**
Platelet lysates containing 20 μg of protein were used. The reaction for assay of TXAS activity was initiated by the addition of TXAS substrate PGH₂ and allowed to proceed for 1 min at 37 °C. The reaction was terminated by the addition of 1 M citric acid, then was neutralized with 1 N NaOH. The concentration of TXA₂ was determined as TXB₂, a stable metabolite of TXA₂, which was measured with a Synergy HT multi-model microplate reader (BioTek Instruments, Winooski, VT, USA) using TXB₂ EIA kit.

**Ex vivo determination of serotonin release**
To investigate the effect on serotonin release, the aggregation was centrifuged at 4 °C for 10 min at 200 × g. The supernatants were used for the assay of serotonin release. Serotonin release was measured with a Synergy HT multi-model microplate reader (BioTek Instruments, Winooski, VT, USA) using serotonin ELISA kit.

**Ex vivo measurement of PT and APTT**
To investigate whether WIB-801CE shows anticoagulant characteristics, if any, has bleeding risk as side effect of anticoagulant [21], we measured PT and APTT, markers of blood coagulation. The PPP (100 μL) was preincubated...
in a two-channel coagulator (Behnk Elektronik GmbH & Co., KG, Norderstedt, Germany) cup (catalog number 95-662, BioMérieux, Marcyl’Etoile, France) with gentle stirring for 1 min at 37 °C. PT was determined as the time interval between the addition of PT reagent (100 μL) to the PPP and the formation of a fibrin clot. After preincubation of PPP for APTT measurement, 100 μL of APTT reagent was added to the PPP (100 μL) and incubated for 3 min at 37 °C. Following incubation, 100 μL of 25 mM CaCl₂ was immediately added to the PPP containing APTT reagent. APTT was determined as the time required to form a fibrin clot.

In vivo tail bleeding time assay
We investigated whether WIB-801CE has bleeding risk, the side effect of antiplatelet substance [21]. WIB-801CE (200, 400 mg/kg-BW) and aspirin (100 mg/kg-BW), a positive control, were orally administered to mice once a day for five days. In this study, we used mice for measuring tail bleeding time according to the method of Kim and Lee [22]. After 5 min of the respective final administration, mice were anesthetized with zoletil (40 mg/kg, i.p.). The distal 0.5 cm segment of the tail was transected with operating knife, and immediately immersed in a tube containing 37 °C of saline. Tail bleeding time was determined as the time required to cause blood coagulation.

In vivo evaluation of anti-acute pulmonary thromboembolism
To confirm the endogenous antithrombotic effect, we used a mice model to generate acute pulmonary thromboembolism [23]. Mice were orally administered with WIB-801CE (200, 400 mg/kg-BW), and aspirin (100 mg/kg-BW). After 1 h of respective administration, the mixture (100 μL) of collagen (300 μg/kg-BW) plus epinephrine (30 μg/kg-BW) were injected via tail vein, and the rate of protection and mortality was observed for 15 min, which were calculated as follow: 1) Protection rate (%) = [(Number of tested mice – Number of dead mice)/Number of tested mice] × 100. 2) Mortality rate (%) = [Number of dead mice/Number of tested mice] × 100. These experiments were proved by the Ethics Committee for animal experiments of Whanin Pharmaceutical Corporation (Suwon, Korea/15-NE-009).

In vitro assay of platelet-mediated fibrin clot retraction
We investigated whether WIB-801CE or cordycepin inhibits fibrin clot retraction, an index of thrombi formation [24]. Human PRP 250 μL (10⁸ platelets/mL) were transferred into polyethylene tube to avoid clot adhesion, then were preincubated with or without WIB-801CE or cordycepin for 10 min at 37 °C, and subsequently stimulated with thrombin (0.5 U/mL) for 60 min at 37 °C. Pictures of fibrin clot were taken at 0 and 60 min using a digital camera, and its quantification was carried out by measurement of clot area using the NIH Image J Software (V1.46, National Institutes of Health, USA). Percentage of clot retraction was calculated as follows: Retraction (%) = [1 - (final clot area/initial clot area)] × 100.

In vitro nitric oxide assay
To observe antiinflammatory effect of WIB-801CE, we used mouse leukemic macrophage RAW264.7 cells. RAW264.7 cells were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA), and were maintained at 37 °C in 5% CO₂ and 95% air in Dulbecco’s Modified Eagle’s Medium (GE Healthcare, Marlborough, USA) containing 10% fetal bovine serum, and 1% penicillin-streptomycin solution. RAW264.7 cells (5 × 10⁴ cells) were preincubated for 30 min with or without WIB-801CE, or inducible nitric oxide synthase (iNOS) inhibitor amino guanidine (AG), and stimulated for 24 h by lipopolysaccharide (10 ng/mL). The supernatant was used for NO assay using Griess reagent. Equal volume of culture supernatant (80 μL) and Griess reagent (80 μL) were mixed. The absorbance of the mixture was measured at 540 nm using spectrophotometer (Spectramax 190, Molecular devices, LLC., Sunnyvale, CA, USA). Nitrite was used as standard of NO.

Ex vivo nitric oxide assay
To investigate the NO production, we used collagen- and ADP-stimulated PRP obtained from rats administered with the WIB-801CE (200, 400 mg/kg-BW). The PRP was centrifuged at 4 °C for 10 min at 10,000 × g to get plasma. The plasma was incubated 1 h with 400 μL methanol:diethyl ether (3:1 mixture v/v), and subsequently plasma proteins were precipitated by centrifuging at 4 °C for 10 min at 10,000 × g. The supernatant was used for NO assay using Griess reagent. Equal volume of supernatant (80 μL) and Griess reagent (80 μL) were mixed. After 30 min, absorbance of the mixture was measured at 540 nm using a Synergy HT multi-model microplate reader (BioTek Instruments, Winooski, VT, USA). Nitrite was used as standard of NO.

In vitro determination of antioxidant activity
To obtain antioxidant effect of WIB-801CE, we measured scavenging activity of free radical in DPPH according to the method [25, 26]. DPPH was dissolved in 99% ethanol to make 200 μM of solution. WIB-801CE and antioxidant AC were dissolved in distilled water. Equal volume of test substances and DPPH were mixed at room temperature. After 30 min, the reduction in DPPH absorbance at 517 nm was measured using spectrophotometer (Optizen 2120UV, Mecasys, Korea). The scavenging activity of DPPH radicals by substances was determined using the following equation [26]: Scavenging activity (%) =
WIB-801CE Peak 1 19.988 ± 0.011 2,278.34 ± 13.77 1,006 ± 0.32 6.93 ± 0.02 69.30 ± 0.20
Pure cordycepin 19.980 2,949.35 100.30 - -

Because it is known that *Cordyceps militaris*, a source of WIB-801CE, has cordycepin (Fig. 1a) [28], we analyzed cordycepin of WIB-801CE with HPLC. As shown in Fig. 1b, peak 1 from WIB-801CE was observed at 19.988 min of the retention time, which was almost in accord with the retention time (19.980 min) of pure cordycepin (Fig. 1c). This means that peak 1 is derived from cordycepin in WIB-801CE. The concentration of peak 1 in WIB-801CE corresponding to cordycepin was 69.30 ± 0.20 mg/g-WIB-801CE (about 6.93 ± 0.02%, Table 1). Whole fruiting body myelia of *Cordyceps militaris* is known to contain 0.16% of cordycepin, but whole fruiting body, stroma, and larva of *Cordyceps sinensis* do not contain cordycepin [29]. Therefore, the cordycepin content in WIB-801CE that we used in this study is very higher than those in whole fruiting body myelia of *Cordyceps militaris*, and in whole fruiting body, stroma, and larva of *Cordyceps sinensis*.

### Results

#### Composition of cordycepin in WIB-801CE

Cordycepin content in WIB-801CE was expressed using the following equation: Cordycepin content (%) = [area of peak 1/area of pure cordycepin] × concentration of pure cordycepin/concentration of WIB-801CE × [(100% - % of water content of cordycepin)/100%] × (% of purity of pure cordycepin/100%) × 100%. Water content of pure cordycepin was 8.18%. Purity of pure cordycepin was 98.0%. The data are given as the mean ± standard deviation (n = 3).

| Table 1 Content of cordycepin in WIB-801CE |
|--------------------------------------------|
| Retention time (min) | Area (mAU × s) | Concentration of sample (µg/mL) | Cordycepin content (%) | Content (mg/g-WIB-801CE) |
|----------------------|----------------|---------------------------------|------------------------|-------------------------|
| Pure cordycepin      | 19.980         | 2,949.35                        | 100.30                 | -                       |
| WIB-801CE Peak 1     | 19.988 ± 0.011 | 2,278.34 ± 13.77                | 1,006 ± 0.32           | 6.93 ± 0.02             | 69.30 ± 0.20 |

Ex vivo measurement of cordycepin effect on rat platelet aggregation

This experiment was performed to investigate the effect of cordycepin on ex vivo platelet aggregation. When cordycepin (15 mg/kg per day) was administered orally to the mice for 14 days, antitumor activity was known to observe [27]. Therefore, we selected 5 and 10 mg/kg-BW per day of cordycepin in this experiment as moderate doses for administration. These doses are corresponded to about 36 and 72% of cordycepin in WIB-801CE (200 mg/kg-BW) that inhibited ex vivo rat platelet aggregation. Rats (Sprague-Dawley, male, 200 g) were acclimatized for a week at a temperature of 24 ± 1 °C and humidity of 55 ± 5%. Before oral administration of cordycepin, rats were fasted for 12 h, then were fed with standard pellets diet (Purina Inc., Korea) had free access to water. Cordycepin (5 and 10 mg/kg-BW) was orally administered to the rats one per day for seven days. Cordycepin were dissolved with distilled water. The experiments were proved by the Ethics Committee for Animal Experiments of Whanin Pharmaceutical Corporation (Suwon, Korea/15-NE-016 for rats). After the final respective administration, all rats were fasted for 24 h, then were anesthetized with 20% urethane before sacrifice. PRP preparation, platelet aggregation, measurement were performed as described before.

#### Protein assay

To determine COX-1, TXAS activity, and protein phosphorylation, protein concentration was measured using bicinchoninic acid assay kit (Pierce Biotechnology, USA).

#### Statistical analyses

The experimental results are indicated as the mean ± standard deviation accompanied by the number of observations. Data were determined by analysis of variance (ANOVA). If this analysis showed significant differences among the group means, then each group was compared by the Newman-Keuls method. Statistical analysis was carried out according to the SPSS 21.0.0.0 (SPSS, Chicago, IL, USA). *p* < 0.05 was considered to be statistically significant.
In vitro effects of WIB-801CE and cordycepin on activation of resting human platelets

Platelet activation is an index of platelet shape change, platelet aggregation and granule secretion, and is the cause of cardiovascular and cerebrovascular disease, and atherosclerosis [33–35]. Accordingly, if WIB-801CE activates resting platelets, unstimulated platelets, a question to evaluate the antiplatelet effects of WIB-801CE might be raised. Therefore, the effect of WIB-801CE and cordycepin on platelet activation was determined by measuring platelet aggregation in resting human platelets. As the results, a positive control collagen (10 μg/mL) activated platelets by increasing platelet aggregation up to 83.3 ± 3.1% (Fig. 1e). However, WIB-801CE (200, 400 μg/mL) alone, and cordycepin (56, 112 μM) alone did not increased platelet aggregation (Fig. 1e), as compared with that (1.0 ± 1.0%) by resting platelets. It was evidenced that WIB-801CE and cordycepin alone do not affect the activation of resting human platelets.

Ex vivo effects of WIB-801CE on platelet aggregation and TXA₂ production

It is known that the inhibition of collagen- and ADP-induced platelet aggregation is potential target to develop antithrombotic agent having antiplatelet characteristics [36, 37]. Therefore, we used on collagen and ADP as agonists. When PRP (10⁶/mL) from control was activated with collagen and ADP, the aggregation rate was increased up to 82.9 ± 6.6% by collagen (Fig. 2a) and 78.7 ± 4.9% by ADP (Fig. 2b). However, collagen- and ADP-induced rat platelet aggregation was significantly attenuated by WIB-801CE (200, 400 mg/kg-BW) (Fig. 2a and b). The inhibitory degrees by 200 mg/kg-BW were 13.0% to that by collagen (Fig. 2a) and 10.9% against that by ADP (Fig. 2b).

We investigated next whether WIB-801CE involves in inhibition of TXA₂ production to attenuate collagen- and ADP-induced rat platelet aggregation. When PRP (10⁶/mL) from control was activated by collagen and ADP, the TXA₂ (determined as TXB₂, a stable metabolite of TXA₂) was increased to 615.0 ± 11.5 ng/10⁸ platelets (Fig. 2c) by collagen and 11.5 ± 1.6 ng/10⁸ platelets by ADP (Fig. 2d). However, collagen- and ADP-produced TXA₂ were inhibited by WIB-801CE (Fig. 2c and d). The inhibitory degrees by 200 mg/kg-BW were 30.7% against that by collagen (Fig. 2c) and 37.3% to that by ADP (Fig. 2d). WIB-801CE (400 mg/kg-BW) inhibited weakly ADP-induced platelet aggregation.
when PRP (10⁸/mL) from WIB-801CE (200, 400 mg/kg-BW) or AA utilization enzymes (COX-1, TXAS). When compared with those by WIB-801CE (200 mg/kg-BW).

ADP-induced AA release (Fig. 3d) and TXAS activity (Fig. 3g) was also attenuated to 27.1% (4.3 ± 0.5 ng/protein-mg/min) by WIB-801CE (200 mg/kg-BW) (Fig. 3f). ADP-induced TXAS activity (5.9 ± 0.5 ng/protein-mg/min) was also attenuated to 27.1% (4.3 ± 0.5 ng/protein-mg/min) by WIB-801CE (200 mg/kg-BW) (Fig. 3g). WIB-801CE (400 mg/kg-BW) inhibited weakly ADP-induced AA release (Fig. 3d) and TXAS activity (Fig. 3g) as compared with those by WIB-801CE (200 mg/kg-BW).

**Ex vivo effects of WIB-801CE on serotonin release**

Next, we investigated the effect of WIB-801CE on serotonin release as an index of granule secretion. Collagen and ADP elevated serotonin release up to 224.5 ± 4.3 ng/10⁸ platelets (Fig. 4a) and 290.1 ± 9.6 ng/10⁸ platelets (Fig. 4b), respectively. On the contrary, collagen- and ADP-released serotonin levels were reduced by WIB-801CE (200, 400 mg/kg-BW) (Fig. 4a and b). The inhibitory degrees by 200 mg/kg-BW were 66.3% to that by collagen (Fig. 4a) and 60.2% against that by ADP (Fig. 4b). WIB-801CE (400 mg/kg-BW) weakly inhibited ADP-induced serotonin release (Fig. 4b) as compared with that by WIB-801CE (200 mg/kg-BW).

**Ex vivo effects of WIB-801CE on p³⁸ MAPK- and ERK-phosphorylation**

It is well reviewed that platelets contain MAPKs such as p³⁸, ERK (1/2), and c-Jun N-terminal kinase [38]. These are activated by various agonists (i.e. collagen, ADP, thrombin, von Willebrand factor, fibrinogen) and subsequently stimulate various enzymes [myosin light chain kinase (MLCK), DG-lipase, cPLA₂, COX-1] associated with platelet activation [9, 10, 38–46]. Of MAPKs, p³⁸ MAPK and ERK2 stimulate MLCK [38–41] to release serotonin, DG-lipase [45] and cPLA₂ [9, 10, 38, 42, 43] to produce AA, TXA₂ precursor. In this study, to observe the relationship with the results that WIB-801CE inhibited collagen- and ADP-induced TXA₂ production (Fig. 2c and d), AA release (Fig. 3c and d) and serotonin release (Fig. 4a and b), we investigated whether WIB-801CE inhibits the phosphorylation of p³⁸ MAPK and ERK.

Collagen and ADP increased potently p³⁸ MAPK phosphorylation (Fig. 5a and b, lane 2) as compared with those by resting platelets (Fig. 5a and b, lane 1), respectively. However, these were diminished by WIB-801CE (200, 400 mg/kg-BW) (Fig. 5a and b, lane 3, 4). WIB-801CE (200 mg/kg-BW) inhibited p³⁸ MAPK phosphorylation up to 52.6% against that by collagen (Fig. 5a, lane 3) and 80.0% against that by ADP (Fig. 5b, lane 3).

Collagen and ADP elevated potently ERK2 (42 kDa) phosphorylation (Fig. 5c and d, lane 2) as compared with those by resting platelets (Fig. 5c and d, lane 1), respectively. However, these were also diminished by WIB-801CE (200, 400 mg/kg-BW) (Fig. 5c and d, lane 3, 4). WIB-801CE (200 mg/kg-BW) inhibited ERK2 (42 kDa) phosphorylation up to 62.5% against that by collagen (Fig. 5c, lane 3) and 81.8% against that by ADP (Fig. 5d, lane 3).

**Ex vivo and in vivo effects of WIB-801CE on blood coagulation and tail bleeding time**

Bleeding is connected to the attenuation of platelet aggregation and blood coagulation, and the inhibition of thrombosis [8, 21, 47, 48]. Accordingly, we investigated the effects...
**Fig. 3** (See legend on next page.)

(a) Western blot analysis of phospholipase A2 (p-PLA$_2$), phospholipase C (p-PLC$_{50}$, p-PLC$_{53}$), phorbol ester (p-PLC$_2$), and β-actin. Collagen (10 μg/mL) and WIB-801CE (mg/kg-BW) concentrations are indicated.

(b) Western blot analysis of phospholipase A2 (p-PLA$_2$), phospholipase C (p-PLC$_{50}$, p-PLC$_{53}$), phorbol ester (p-PLC$_2$), and β-actin. ADP (5 μM) and WIB-801CE (mg/kg-BW) concentrations are indicated.

(c) Graph showing the release of arachidonic acid in response to Collagen (10 μg/mL) and WIB-801CE (mg/kg-BW) treatments. Significant differences are indicated with asterisks (*p < 0.05).

(d) Graph showing the release of arachidonic acid in response to ADP (5 μM) and WIB-801CE (mg/kg-BW) treatments. Significant differences are indicated with asterisks (*p < 0.05).

(e) Graph showing the COX1 activity in response to Collagen (10 μg/mL) and WIB-801CE (mg/kg-BW) treatments. Non-significant differences are indicated with ‘NS’.

(f) Graph showing the TxA$_3$ activity in response to Collagen (10 μg/mL) and WIB-801CE (mg/kg-BW) treatments. Significant differences are indicated with asterisks (*p < 0.05).

(g) Graph showing the TxA$_3$ activity in response to ADP (5 μM) and WIB-801CE (mg/kg-BW) treatments. Significant differences are indicated with asterisks (*p < 0.05).
of WIB-801CE on ex vivo blood coagulation time (PT, APTT) and in vivo tail bleeding time as indexes of bleeding. As shown in Table 2, both ex vivo PT and APTT were not significantly prolonged by PPP from WIB-801CE rats as compared with those (PT, 13.2 ± 2.3 s; APTT, 26.2 ± 4.1 s) by normal. Warfarin (1 mg/kg-BW) infinitely prolonged PT and APTT (Table 2).

With regard to the effects of WIB-801CE on tail bleeding time, WIB-801CE (200, 400 mg/kg-BW) significantly prolonged from 125.3 ± 17.0 s by control to 264.8 ± 79.0 s and 360.3 ± 83.8 s, respectively (Table 3). Aspirin (100 mg/kg-BW) also prolonged tail bleeding time to 1,800.0 ± 0.0 s (Table 3). Aspirin (100 mg/kg-BW) potently prolonged bleeding time to 1,336.5%, on the other hand, WIB-801CE (200, 400 mg/kg-BW) prolonged it up to 111.4% and 187.5% as compared with that (125.3 ± 17.0 s) by control, respectively (Table 3).

In vivo effects of WIB-801CE on acute pulmonary thromboembolism

Because antiplatelet drugs play an important role in protection of thrombus formation, we investigated whether WIB-801CE, inhibiting ex vivo platelet aggregation, has also a protective effect on endogenous thrombus formation. In this study, in vivo venous antithrombotic effect of WIB-801CE was estimated using collagen plus epinephrine-induced acute pulmonary thromboembolism mouse model [23, 49–51]. As shown in Table 4, when the mixture of collagen plus epinephrine was treated to mice, the protection rate was 4.2% against acute pulmonary thromboembolism, and the mortality rate was 95.8%. However, in WIB-801CE-treated mice, the protection degree from a pulmonary thromboembolism was increased to 25.0% by WIB-801CE (200 mg/kg-BW), and 35.0% by WIB-801CE (400 mg/kg-BW) in a dose dependent manner (Table 4). In aspirin (100 mg/kg-BW)-treated mice, the protection degree from a pulmonary thromboembolism was increased up to 35.0%, and the mortality was decreased to 65.0%, which were equal to those by WIB-801CE (400 mg/kg-BW)-treatment (Table 4). These mean that WIB-801CE is actually valid to protect venous thromboembolism like aspirin.

In vitro effects of WIB-801CE and cordycepin on fibrin clot retraction

Fibrin clot retraction is a final index of platelet aggregation-mediated thrombotic formation, and is resulted from interaction of fibrin-platelet [24]. Thrombin stimulated the retraction of fibrin clot (Fig. 6a, dotted circle), but WIB-801CE (200, 400 μg/mL) inhibited it in a dose dependent manner (Fig. 6a and b). WIB-801CE (200 μg/mL) inhibited
thrombin-induced fibrin clot retraction up to 74.3% (Fig. 6b). Cordycepin (56, 112 μM) corresponding to dose (200, 400 μg/mL) of WIB-801CE attenuated thrombin-induced fibrin clot retraction in a dose dependent manner (Fig. 6a and b). Cordycepin (56 μM) corresponding to 200 μg/mL of WIB-801CE inhibited thrombin-induced fibrin clot retraction (14.4 ± 5.1%) up to 59.1% (Fig. 6b).

**In vitro and ex vivo effects of WIB-801CE on NO production**

It is well known that monocytes/macrophages and neutrophils produce various inflammatory mediators (i.e. NO, Table 2 Ex vivo effects of WIB-801CE-administration on blood coagulation

| Animal group | Dose (mg/kg-BW) | PT (s) | APTT (s) | N  |
|--------------|-----------------|--------|-----------|----|
| Normal       | N.D.            | 13.2 ± 2.3 | 26.2 ± 4.1 | 8  |
| WIB-801CE    | 200             | 17.7 ± 4.9S | 26.3 ± 3.9S | 8  |
|              | 400             | 16.8 ± 3.2S | 26.0 ± 3.9S | 8  |
| Warfarin     | 1               | ∞       | ∞         | 6  |

The results were expressed as the mean ± standard deviation (n = 8 or 6). ∞, no coagulation; NS not significant versus normal; N.D normal diet. N number of tested rats; BW body weight, PT prothrombin time, APTT activated partial thromboplastin time, s, second.
prostaglandin E₂), and subsequently activate platelets to generate atherothrombosis [52, 53]. In recent, it is also reported that neutrophil-produced NO activates platelet in chronic renal failure [54]. Accordingly, we investigated whether WIB-801CE inhibits in vitro NO production in RAW264.7 macrophage cells. As shown in Fig. 7a, lipo-polysaccharide (LPS), an activator of macrophages, potently produced NO as compared with that of normal. However, WIB-801CE dose (15 to 50 μg/mL)-dependently attenuated LPS-elevated NO production (Fig. 7a). iNOS inhibitor AG potently inhibited NO production (Fig. 7a).

The ex vivo NO levels in plasma from PRP of control (Fig. 7b, lane 2, 5) and WIB-801CE (200, 400 mg/kg-BW) stimulated with collagen (Fig. 7b, lane 3, 4) and ADP (Fig. 7b, lane 6, 7) were not changed as compared with that (Fig. 7b, lane 1) in plasma from control rat, in the absence of ADP, collagen, and WIB-801CE. These mean that WIB-801CE (200, 400 mg/kg-BW) did not at least activate inflammatory iNOS in leukocytes (i.e. monocyte, macrophage, neutrophil) in vivo. Otherwise, the ex vivo NO levels would be increased in plasma from WIB-801CE (200, 400 mg/kg-BW) stimulated with collagen and ADP.

Ex vivo effects of cordycepin on platelet aggregation
As shown in Table 5, When PRP (10⁸/mL) from control, was activated with ADP (10 μM), the aggregation rate was increased up to 75.7 ± 0.6%. But ADP-induced platelet aggregation significantly decreased by cordycepin (5, 10 mg/kg-BW) (Table 5). The inhibitory degree by 5 mg/kg-BW was 31.7%. This is higher than that (10.9%) by WIB-801CE (200 mg/kg-BW) (Fig. 2b).

Discussion
WIB-801CE and its component cordycepin did not affect the cytotoxicity (determined as LDH leakage) and platelet activation (determined as platelet aggregation) to resting human platelets in vitro. This means that there is no problem to evaluate the antiplatelet effects of WIB-801CE ex vivo. It is well established that various agonists (i.e. collagen, ADP, thrombin)-produced TXA₂ generates circulatory disorder such as thrombosis, atherosclerosis, and myocardial infarction by stimulating platelet aggregation, vasoconstriction, and bronchoconstriction [5, 58, 59]. Therefore, it is essential to inhibit platelet aggregation and TXA₂ production to prevent circulatory disorder in blood vessel. WIB-801CE attenuated ex vivo collagen- and ADP-induced platelet aggregation and TXA₂ production.

These results were connected to the ex vivo inhibition of AA release and TXAS activity by WIB-801CE in collagen- and ADP-activated platelets. WIB-801CE did not inhibit ex vivo collagen- and ADP-induced PLCβ₃ (Ser⁵³⁷, Ser¹¹⁰⁵), PLCγ₂ (Tyr¹²¹⁷) phosphorylation. These mean that WIB-801CE would produce DG from phosphatidylinositol 4,5-bisphosphate in collagen- and ADP-activated platelets. DG is known to hydrolyze by p³⁸ MAPK-activated DG-lipase to release AA [45]. If so, it is considered that WIB-801CE may attenuate AA release by inhibiting p³⁸ MAPK/DG-lipase pathway [45] without affecting inhibition of cPLA₂ and PLCβ₃ [9, 10, 42, 43]. Because agonist-produced TXA₂ enforces thrombus formation as a positive promoter [5, 58, 59], a compound or substance that inhibits the activity of COX-1 or TXAS, the production of TXA₂ or the action of

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**Table 3** In vivo effects of WIB-801CE on tail bleeding time

| Animal group | Dose (mg/kg-BW) | Tail bleeding time (s) | Δ (%) | N |
|--------------|-----------------|-----------------------|------|---|
| Control      | Saline          | 125.3 ± 17.0          | 0    | 8 |
| WIB-801CE    | 200             | 264.8 ± 79.0          | 111.3| 8 |
|              | 400             | 360.3 ± 83.8**        | 187.5| 8 |
| Aspirin      | 100             | 1,800.0 ± 0.0**       | 1,336.5| 5 |

The results were expressed as the mean ± standard deviation (n = 8 or 5). Δp < 0.05 compared with control. **p < 0.001 compared with control. N number of tested mice, BW body weight, Δ second Δ (%) = (WIB-801CE or aspirin)/control × control × 100.

**Table 4** In vivo effects of WIB-801CE on acute pulmonary thromboembolism

| Animal group | Dose (mg/kg-BW) | No. of tested mice¹⁵ | No. of dead¹⁶ | Protection (%) | Mortality (%) |
|--------------|-----------------|----------------------|--------------|----------------|--------------|
| Control      | -               | 24                   | 23           | 4.2            | 95.8         |
| WIB-801CE    | 200             | 20                   | 15           | 25.0           | 75.0         |
|              | 400             | 20                   | 13           | 35.0           | 65.0         |
| Aspirin      | 100             | 20                   | 13           | 35.0           | 65.0         |

Protection (%) = (ΔO - ΔI)/ΔI × 100, Mortality (%) = ΔI/O × 100. No., number of tested mice; BW, body weight.
TXA₂ is evaluated as antithrombotic agents. Many studies have been performed to discover therapeutic agents that can counteract the effects of TXA₂. Various phytochemicals (i.e. epigallocatechin-3-gallate, caffeic acid, chlorogenic acid, caffeidine, sanguinarine) are known to inhibit COX-1 rather than TXAS to suppress the TXA₂ production in vitro or ex vivo [60–65]. However, WIB-801CE inhibited the activity of TXAS rather than COX-1 ex vivo, which reflects that WIB-801CE inhibits the TXA₂ production pathway from PGH₂ rather than prostaglandin G₂ production pathway from AA. In this study, we showed that WIB-801CE may involve in down-regulation of both p³θ MAPK phosphorylation to vanish the AA supply from DG and TXAS activity to block the TXA₂ production from PGH₂ ex vivo. Therefore, it is apparent that WIB-801CE can be beneficially used to prevent the TXA₂-mediated thrombus formation in vivo.

It is well known that agonist-released serotonin stimulates irreversibly platelet aggregation, and subsequently causes the thrombosis as well as TXA₂ [66–68]. WIB-801CE inhibited collagen- and ADP-induced serotonin release ex vivo, which reflects that WIB-801CE can inhibit the irreversible platelet aggregation in vivo. WIB-801CE potently inhibited ex vivo the phosphorylation of p³θ MAPK and ERK2 (42 kDa), but not Ca²⁺-dependent
Fig. 7 (See legend on next page.)
myosin light chain (MLC) phosphorylation (Data not shown) that involves in serotonin release [14, 69–72] in collagen- and ADP-activated platelets. These results are allowed to consider that WIB-801CE seems to attenuate ex vivo serotonin release by inhibiting the phosphorylation of p38 MAPK and ERK2 rather than MLC in collagen- and ADP-activated platelets. This is similar to the reports that some phytochemicals (i.e. caffeic acid phenethyl ester, ginsenoside Rp1) inhibit collagen- and ADP-induced ATP release by phosphorylating p38 MAPK and ERK2 [39, 73]. In recent, we found that CE-WIB801C, n-butanol extracts from Cordyceps militaris, and cordycepin purified from CE-WIB801C has in vitro antiplatelet effect by inhibiting fibrinogen binding to glycoprotein IIb/IIIa via stimulation of CAMP-dependent phosphorylation of vasodilator-stimulated phosphoprotein (Ser157), and inhibition of phosphatidylinositol-3 kinase/Akt phosphorylation [74]. Antiplatelet effect of CE-WIB801C was involved in inhibition of collagen-induced serotonin release. Accordingly, we confirmed that the extracts from Cordyceps militaris have antiplatelet effect in vitro and in vivo.

The formation of fibrin clot by intrinsic and extrinsic blood coagulation factors together with platelet aggregation at injured blood vessels is another cause of thrombogenesis. WIB-801CE did not significantly prolong PT and APTT as compared with that by normal ex vivo. This reflects that WIB-801CE has no anticoagulant characteristics. WIB-801CE, however, weakly prolonged average PT as compared with that by normal. The prolongation of PT is associated with the reduction of coagulation factor VII production by inhibition of NADPH-vitamin K reductase [75]. The dose (1 mg/kg-BW) of warfarin, an inhibitor of NADPH-vitamin K reductase, that unlimitedly prolonged PT is corresponded to high dose (60 mg/day) in case of giving to human (60 kg), which is more 12 fold than international normalized dose (5 mg/day) of warfarin [76]. In this study, because it is unknown whether the weak extension of average PT by WIB-801CE is clinically safety or risk, it is necessary to investigate PT using international normalized dose (5 mg/day) of warfarin, then the safety of WIB-801CE in the weak extension of PT should be evaluated in the future.

In addition, WIB-801CE without significantly affecting the prolongation of blood coagulation time may not influence on inhibition of fibrin production. If so, because the fibrin is retracted by the interaction with platelet aggregation [77], anybody may apprehend that the thrombus could be generated by WIB-801CE in vivo. But it is considered that its fear can be excluded because WIB-801CE inhibited both thrombogenic TXA2 production and serotonin release ex vivo, and thrombin-induced fibrin clot retraction in vitro. These results mean that WIB-801CE can strongly inhibit the fibrin clot retraction by down-regulating platelet activation without significantly affecting the blood coagulation. This is also evidenced as the effect that WIB-801CE inhibited collagen plus epinephrine-induced acute pulmonary thromboembolism in vivo, which is a marker of platelet aggregation-generated thrombogenesis. At the present study, however, it is unknown whether cordycepin in WIB-801CE contributed to the inhibition of acute pulmonary thromboembolism. These should be studied in the future.

As well as anticoagulants, antiplatelet drugs (i.e. aspirin, clopidogrel) also cause bleeding, and surprisingly may generate blood loss [21]. It is known that 20–40 mg/day of aspirin is clinically used in human to protect thrombotic disease [78]. The dose (100 mg/kg-BW) of aspirin, a positive control, seriously prolonged tail bleeding time of mice as compared with that by control. This aspirin dose (100 mg/kg-BW) is corresponded to high dose (6,000 mg/day) in case of giving to human (60 kg), which is more 300-150 fold than clinical dose (20–40 mg/day) of aspirin and impossible to compare with WIB-801CE in tail bleeding time of mice. At the present study, because it is unknown whether the significant extension of tail bleeding time by WIB-801CE is clinically safety or risk, it is necessary to investigate tail bleeding time using the clinical dose (20–40 mg/day) of aspirin as positive control, then the safety of WIB-801CE-prolonged tail bleeding time should be evaluated in the future.

Leukocyte-produced ROS oxidizes low density lipoprotein (LDL) in blood, then oxidized LDL (ox-LDL) is incorporated into macrophage to generate foam cell

Table 5 Effects of cordoncycin administration in ADP-induced rat platelet aggregation

|                  | Light transmission (%) | Inhibition (%) | Inhibition (%) |
|------------------|------------------------|---------------|---------------|
| Control + ADP (10 μM) | 75.7 ± 0.6             | -             |    |
| Cordycepin (5 mg/kg-BW) + ADP (10 μM) | 51.7 ± 0.6** | 31.7          |    |
| Cordycepin (10 mg/kg-BW) + ADP (10 μM) | 47.3 ± 2.5** | 37.4          |    |

The data are given as the mean ± standard deviation (n = 3). **p < 0.001 compared with control. Inhibition (%) = [(cordycepin + ADP) – control]/control × 100 (%)
which damages vascular wall by inducing inflammation. Because platelet aggregation is caused at injured place of vascular wall, the inflammation by leukocyte-produced ROS and NO is the cause of thrombus formation. This means that the counteraction of agonists-induced platelet aggregation and leukocyte-induced inflammation might be contributed to the inhibition of thrombosis. WIB-801CE had ex vivo inhibitory effects on the release of serotonin that stimulates the uptake of ox-LDL into macrophage [79]. In addition, WIB-801CE inhibited NO production and elevated scavenging activity of free radical in DPPH in vitro. Therefore, it is anticipated that WIB-801CE would not activate inflammatory leukocytes in vivo as evidenced that WIB-801CE dose not affect ex vivo NO production. These results suggest that WIB-801CE might have antithrombotic effects by inhibiting inflammation via antioxidative action.

Because we could not identify cordycepin or its metabolites in PRP from WIB-801CE (200, 400 mg/kg-BW) rats, in the present study, we could not explain whether cordycepin in WIB-801CE was absorbed through intestine and subsequently involved in inhibition of platelet aggregation. This study should be performed in the future. Considering antiplatelet effects observed by blood collection of 2 h after final administration of WIB-801CE and 14 days after administration of cordycepin, although cordycepin is metabolized to an inactive 3′-deoxyhypoxanthosine by adenosine deaminase in rat blood [80–82], it could be thought that unknown substances in WIB-801CE or cordycepin-derived unknown substance might involve in inhibition of platelet aggregation in an acute or chronic manner.

WIB-801CE (200, 400 mg/kg-BW)-dose independently exerted its inhibitory effects on platelet aggregation, TXA2 production, AA release, TXAS activity, serotonin release, p38 MAPK phosphorylation and ERK2 phosphorylation in ADP-activated platelets. Considering the inhibition of ADP-induced platelet aggregation is a potential target to develop antithrombotic agent having antiplatelet characteristics [36, 37], it is thought that high dose of WIB-801CE (400 mg/kg-BW) might exert undesirable effect on platelets in vivo.

Conclusion

Cordycepin-enriched WIB-801CE from Cordyceps militaris vanished ex vivo thrombogenic molecules (TXA2, serotonin), and their associated signaling molecules (AA, TXAS, p38 MAPK, ERK2) in platelet aggregation. Furthermore, WIB-801CE inhibited in vivo acute pulmonary thromboembolism, an index of thrombosis formation, without having cytotoxicity and risk of serious bleeding, but with antioxidant and antiinflammatory activity. Therefore, we suggest that WIB-801CE may be a beneficial and effective substance to treat or protect thrombosis, atherosclerosis, and myocardial infarction via inhibition of platelet activation in vivo.

Abbreviations

AA: Arachidonic acid; AC: Ascorbic acid; ADP: Adenosine diphosphate; AG: Amino guanidine; ANOVA: Analysis of variance; APTT: Activated partial thromboplastin time; ATCC: American type culture collection; BW: Body weight; COX-1: Cyclooxygenase-1; cPLA2: Cytosolic phospholipase A2; DG: Diacylglycerol; DPPH: 2,2-diphenyl-1-picrylhydrazyl; ECL: Enhanced chemiluminescence solution; IA: Enzyme immunsassay; ELSA: Enzyme-linked immunosorbent assay; ERK: Extracellular signal-regulated kinase; HPLC: High performance liquid chromatography; HRP: Horseradish peroxidase conjugate; ICR: Institute of Cancer Research; IgG: Immunoglobulin G; INOS: Inducible nitric oxide synthase; KBRC: Korean Red Cross Blood Center; LDH: Lactate dehydrogenase; LDL: Low density lipoprotein; LPS: Lypopolysaccharide; MAPK: Mitogen-activated protein kinase; ML: Myosin light chain; MLCK: Myosin light chain kinase; NO: Nitric oxide; ox-LDL: Oxidized low density lipoprotein; PGH2: Prostaglandin H2; PLC: Phospholipase C; PLCγ: Phospholipase Cγ; PPP: Platelet-poor plasma; PRP: Platelet-rich plasma; PT: Prothrombin time; PVDF: Polyvinylidene difluoride; ROS: Reactive oxygen species; SDS-PAGE: Sodium dodecyl sulfate-polyacrylamide gel electrophoresis; TXA2: Thromboxane A2; TXAS: TXA2 synthase; TXB2: Thromboxane B2; WIB-801CE: (Compound from 2008 First Project of Bioteam, Whanin Pharm. Co., Ltd., Suwon, Korea), an ethanol extract from Cordyceps militaris-hypha

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Availability of data and materials

The contents of this manuscript was argued and are useful on requirement from the corresponding author.

Authors’ contributions

Conception and design of the experiment: HJP, ILK. Performance of the experiment: HWK, JHS, DHL, WJO, GSN, MJK, HKH, JYN, JYL. Analysis and arrangement of the data: HWK, JHS, DHL, WJO, GSN, MJK, HKH, JYN, JYL, HHK. Contribution of reagents, materials, and tool: HWK, JHS, DHL. Interpretation of the data: HWK, JHS, DHL, WJO, HJP. Writing of initial paper drafts: HJP. Contribution of manuscript preparation: HWK, JHS, DHL, HHK. All authors read and approved the final manuscript.

Competing interests

Inje University has performed the measurement of cytotoxicity, platelet aggregation, TXA2, activities of cPLA2, PLCγ, and PLCβ, arachidonic acid release, serotonin release, COX-1 activity, TXAS activity, phosphorylation of p38 MAPK, and ERKs, fibrin clot retraction, NO production of WIB-801CE (200, 400 mg/kg-BW), oxidation, ex vivo antiplatelet effect of cordycepin and supplemental experiments. These above experiments are supported by the Basic Science Research Program via the National Research Foundation of Korea (NRF) funded by the Ministry of Education, Science and Technology, Korea (NRF-2011-0012143 to Hwa-Jin Park). Authors of Inje University declare that they have no potential competing interests.

Consent for publication

The authors agreed to publish the results of this study.

Ethics approval and consent to participate

The experiments of human platelets were followed the guidelines of The Korea National Institute for Bioethics Policy Public Institutional Review Board (Seoul, Korea/PBR12-072-01). The animal experiments were performed as per the Ethics Committee of Whanin Pharmaceutical Corporation (Suwon, Korea/
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