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OPEN Cannabinoid receptor subtype 2 (CB₂R) agonist, GW405833 reduces agonist-induced Ca²⁺ oscillations in mouse pancreatic acinar cells

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Emerging evidence demonstrates that the blockade of intracellular Ca²⁺ signals may protect pancreatic acinar cells against Ca²⁺ overload, intracellular protease activation, and necrosis. The activation of cannabinoid receptor subtype 2 (CB₂R) prevents acinar cell pathogenesis in animal models of acute pancreatitis. However, whether CB₂Rs modulate intracellular Ca²⁺ signals in pancreatic acinar cells is largely unknown. We evaluated the roles of CB₂R agonist, GW405833 (GW) in agonist-induced Ca²⁺ oscillations in pancreatic acinar cells using multiple experimental approaches with acute dissociated pancreatic acinar cells prepared from wild type, CB₁R-knockout (KO), and CB₂R-KO mice. Immunohistochemical labeling revealed that CB₂R protein was expressed in mouse pancreatic acinar cells. Electrophysiological experiments showed that activation of CB₂Rs by GW reduced acetylcholine (ACh)-, but not cholecystokinin (CCK)-induced Ca²⁺ oscillations in a concentration-dependent manner; this inhibition was prevented by a selective CB₂R antagonist, AM630, or was absent in CB₂R-KO but not CB₁R-KO mice. In addition, GW eliminated L-arginine-induced enhancement of Ca²⁺ oscillations, pancreatic amylase, and pulmonary myeloperoxidase. Collectively, we provide novel evidence that activation of CB₂Rs eliminates ACh-induced Ca²⁺ oscillations and L-arginine-induced enhancement of Ca²⁺ signaling in mouse pancreatic acinar cells, which suggests a potential cellular mechanism of CB₂Rmediated protection in acute pancreatitis.

Acute pancreatitis is an inflammatory disease, which has several causes and symptoms and requires immediate medical attention^{1,2}. In clinical practice, there are still no efficient drugs that specifically treat acute pancreatitis¹. Emerging evidence demonstrates that a primary event initiating the process of acute pancreatitis is the excessive release of Ca^{2+} from intracellular stores³. These studies provide a promising therapeutic strategy—the blockade of Ca^{2+} release-activated Ca^{2+} currents in pancreatic acinar cells may provide significant protection against Ca^{2+} overload, intracellular protease activation, and necrosis, which are the major triggers of acute pancreatitis.

The cannabinoid receptor type 2 (CB₂R) is a G protein-coupled receptor that, in humans, is encoded by the CNR2 gene⁴. CB₂Rs are predominantly expressed in the periphery, especially in immune cells, suggesting that CB₂R mediates the effects of cannabinoids mainly in the immune system. For example, activation of CB₂Rs

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inhibits adenylyl cyclase via Gi/Go $_{\alpha}$ subunits and causes a reduction in the intracellular level of cyclic adenosine monophosphate (cAMP)^{5,6}, which has been implicated in a variety of modulatory functions including immune suppression, induction of apoptosis, and induction of cell migration⁷. Thus, CB₂R agonists may be useful candidates for treating inflammatory diseases and pain⁸. Consistent with these findings, increased CB₂R expression has been observed in spinal cord, dorsal root ganglion, and activated microglia in a rodent model of neuropathic pain, as well as in human hepatocellular carcinoma tumor samples⁹. In addition, emerging data demonstrate that CB₂R mRNA and protein are expressed in pancreatic acinar cells, and activation of these CB₂Rs prevents acinar cell pathogenesis in an animal model of pancreatitis¹⁰. However, whether the activation of CB₂R modulates intracellular Ca²⁺ signals in pancreatic acinar cells is largely unknown. Specifically, it is unknown whether an agent that induces pancreatitis (e.g., L-arginine) enhances Ca²⁺ oscillations and whether application of a CB₂R agonist eliminates L-arginine-induced enhancement of Ca²⁺ oscillations in pancreatic acinar cells.

In this study, we address these important questions using patch-clamp and confocal Ca^{2+} imaging approaches combined with immunohistochemistry using wild-type (WT), CB_1R -knockout (KO), and CB_2R -KO mice.

Methods

All experimental protocols were approved by and performed in accordance with guidelines set by the animal care and use and ethical committees at the Barrow Neurological Institute, Xiangya Hospital (Hunan, Changsha, China), and Shantou University Medical College (Shantou, Guangdong, China).

Animals. Mice used for this study were adult (4–6 month old), male, CD1 mice (Charles River Laboratories International, Inc., Wilmington, MA, USA). In addition, WT, CB₁RKO¹¹, and CB₂RKO mice¹² with C57BL/6J genetic backgrounds were initially provided by Dr. Zheng-Xiong Xi at the National Institute on Drug Abuse (NIDA; Bethesda, MD, USA), and were then bred in animal facilities at the Barrow Neurological Institute, which are accredited by the Association for Assessment and Accreditation of Laboratory Animal Care. Genotyping was performed at the NIDA Intramural Research Program before experiments were begun. All animals used in the experiments were matched for age (8–14 weeks) and weight (25–35 grams).

Mouse Pancreatic Acinar Cell Preparation. Acute isolated pancreatic cells were prepared as previously described¹³⁻¹⁵. In brief, pancreatic glands were taken from isoflurane-anesthetized mice, and fragments of the tissue were minced and digested using collagenase (200 U/mL, 25–30 min, 37 °C; Wako Pure Chemicals, Osaka, Japan) in the presence of 1-mM Ca²⁺. After collagenase digestion, the cell suspension was gently pipetted to obtain further separation of the cells, and then washed with physiological saline. A 100-µL volume of cell suspension was then poured into extracellular solution in a 2-mL experimental bath. The isolated cells usually adhered to the bottom within 15–20 min and were used for recording within 3 h after preparation. All experiments were performed at room temperature (22 ± 1 °C).

Whole-Cell Patch-Clamp Recording and Perforated-Patch Recording. Conventional whole-cell patch-clamp recording was used to record the Ca²⁺-activated Cl⁻ currents for monitoring intracellular Ca²⁺ signal oscillations, as reported previously^{13,14}. The recording pipettes, made from borosilicate glass capillaries, had a resistance of 3–5 MΩ when filled with pipette solution. After a GΩ seal was established between the cell membrane and the pipette, a whole-cell configuration was achieved by brief negative suction. Transmembrane currents were recorded with a patch-clamp amplifier (Axopatch 200B; Molecular Devices, Sunnyvale, CA, USA) at a holding potential (V_H) of -30 mV. For perforated-patch recording, amphotericin B (200 µg/mL) was dissolved into the pipette solution. In these studies, we did not compensate for series resistance.

Drug Application. A stream of standard extracellular solution was continuously perfused over the cell during recording. A computer-controlled U-tube system was used for drug application¹⁶. For intracellular drug application, the drug was added into pipette solution, and establishment of a whole-cell configuration allowed the drug to diffuse into the recorded cell.

Amylase Estimation. Serum amylase activity was measured using the AMS assay kit (Nanjing Jiancheng Corp., Nanjing, China) and a microplate reader, following the manufacturer's recommendations.

Myeloperoxidase Estimation. To measure myeloperoxidase (MPO) activity, lung tissues were immediately homogenized on ice in 10 volumes of normal saline. MPO activity was measured using the MPO assay kit (Nanjing Jiancheng Corp., Nanjing, China) and a microplate reader, following the manufacturer's recommendations.

CB₂**R Immunoblot Assay.** WT, CB₁R-KO, and CB₂R- KO mice (3 mice for each group) were anesthetized and quickly perfused with saline to flash all blood cells. Both whole striatum and spleen tissue were dissected out, snap frozen, and kept on dry ice. All the tissues were homogenized in cell lysis buffer (Cell Signaling Technology, Inc., Danvers, MA, USA) using a sonicator and centrifuge at 15,000 rpm for 15 min at 4 °C to get supernatant. The protein concentration for each sample was quantified with a Bio-Rad Protein Assay (Bio-Rad Laboratories, Hercules, CA, USA). A total of 20- μ g protein (spleen) or 40- μ g protein (striatum) were loaded and separated by SDS-PAGE in a 4–15% gradient gel for the detection of endogenous calnexin (Enzo Life Sciences, SPA865) and CB₂R (NIDA-5633) by using Invitrogen blotting and transferring modules (Grand Island, NY, USA). Membranes were blocked for 2 h at room temperature with Licor Odyssey blocking buffer (LI-COR Biosciences, Lincoln, NE, USA) after washing 3 times with phosphate-buffered saline containing 0.1% Tween-20. Membranes were first incubated with either anti-CB₂ (1:500 NIDA-5633 Ab) or anti-calnexin (1:1,000) antibody overnight at 4°C. After washing 3 times, the membranes were incubated with goat anti-rabbit IgG (IRDye 680CW) (1:2,500) for 1.5 h at room temperature. Then the membranes were washed 3 times and then scanned in a Licor Odyssey Sa Imaging System (LI-COR Biosciences).

Immunohistochemistry. Sections were first blocked in 5% bovine serum albumin (BSA) and 0.5% Triton X-100 in phosphate buffer (PB) for 2 h at room temperature. Then, sections were incubated with 1:500 NIDA-5633 mCB₂R antibody (Genemed Synthesis Inc, San Antonio, TX, USA) at 4 °C overnight. After washing 3 times with 0.1 M PB, sections were incubated with Alexa Fluor 488 goat anti-rabbit secondary antibody (Invitrogen, Carlsbad, CA, USA) in 5% BSA and 0.5% Triton X-100 PB for 2 h at room temperature. Sections were then washed, mounted, and cover slipped. Images were taken with a fluorescence microscope (Nikon Eclipse 80i) equipped with a digital camera (Nikon Instruments Inc., Melville, NY, USA).

Confocal Ca²⁺ Imaging. Dissociated pancreatic acinar cells were first incubated with fluo-4-AM (15 μ M) (Molecular Probes, Eugene, OR, USA) for 15 min, followed by a 10-min rest allowing for de-esterification of the indicator. Confocal imaging was performed using an Olympus FluoView FV1000 microscope (Olympus Corporation, Center Valley, PA, USA) equipped with an argon laser (488 nm) and a UPLSAPO 40×, 0.95 NA objective. X-Y imaging was performed at a rate of 1.644 s per frame, 400 frames total, with a resolution of 512 × 512. Fluorescent fluo-4 signal was measured using ImageJ v.1.47 (available from the U.S. National Institutes of Health, Bethesda, MD, USA; http://imagej.nih.gov/ij/).

Solution and Chemicals. Standard extracellular solution contained (in mM): 140 NaCl, 1.0 CaCl₂, 4.7 KCl, 1.13 MgCl₂, 10 glucose, and 10 HEPES, adjusted to pH 7.2 with NaOH. Pipette solution contained (in mM) 140 KCl, 1.13 MgCl₂, 5 Na₂ATP, 0.24 EGTA, 10 glucose and 10 HEPES, pH 7.2. Drugs used in this study were GW405833 (Supplemental Fig. 1), JWH133, ACEA, and AM630, cholecystokinin (CCK), which were purchased from Tocris Bioscience (Minneapolis, MN, USA). Acetylcholine (ACh), amphotericin B, and L-arginine were purchased from Sigma-Aldrich (St. Louis, MO, USA).

CB₂R antibodies, NIDA-5633 mCB2-Ab (customer-designed, raised in rabbit) that recognize the C-terminal (326-340 aa) of mCB₂Rs, were produced by Genemed Synthesis Inc. (San Antonio, TX, USA).

Statistics. For patch-clamp experiments, the Ca^{2+} -activated Cl^- current responses were presented as the current charge (current area/Cm/min), and then the drug-induced changes were compared to the baseline level of charge (induced by ACh). When data were obtained from the same recorded cell and the changes of ACh response were compared before, during, and after testing drug exposure, a paired Student *t* test was used. To compare the effect of the tested drug between 2 groups of animals (e.g., saline group and L-arginine group), the unpaired Student *t* test was used. To analyze multiple effects, one-way analysis of variance (ANOVA) with Tukey's post hoc tests were used.

Results

CB₂**Rs Are Expressed on Mouse Pancreatic Acinar Cells.** Under the acutely dissociated acinar cell protocol, the isolated cells exhibited a typical kidney shape with secretion granules in the central area of the cells (Supplemental Fig. 2), suggesting the purity of the acinar cells as previously reported^{13,15,17}. Figure 1A shows the results of the immunoblot assays, illustrating that a CB₂-positive band was detected (at ~40 kD) in both the spleen and striatal tissues of WT and CB₁-KO (CB₁^{-/-}), while the densities of this band in CB₂-KO mice (CB₂^{-/-}) were substantially reduced in CB₂-rich spleen tissues and almost undetectable in striatal tissues. Figure 1B shows CB₂R immunostaining with the CB₂R antibody (NIDA-5633), illustrating that the high densities of CB₂R immunostaining was detected in a minority of spleen cells of WT mice. In contrast, a very low density of CB₂-like staining was detected in a minority of spleen cells in CB₂R KO mice, suggesting that the NIDA-5633 antibody used is highly mouse CB₂R-specific. We then used this antibody to detect CB₂R expression in single isolated acinar cells. Figure 1C demonstrates the photographs taken in bright field (Ca), mouse CB₂R antibody (mCB₂-ir, Cb), DAPI (Cc), and merged mCB₂-ir and DAPI (Cd). We found high densities of CB₂R immunolabeling in pancreatic acinar cells (Fig. 1Cb,d). These results suggest that CB₂R protein is expressed in dissociated mouse pancreatic acinar cells.

Effects of GW405833 on ACh-Induced Ca²⁺ Oscillations. In acutely dissociated pancreatic acinar cells, low nanomolar concentrations of ACh induced intracellular Ca²⁺ signal oscillations, which can be detected using patch-clamp recording and Ca²⁺ imaging as previously reported^{13,14,18-20}. Our initial series of experiments was designed to test the effects of the CB₂R agonist, GW405833 (GW), on ACh-induced Ca²⁺ oscillations. Figure 2A demonstrates an experimental protocol, in which the ACh (e.g., 10 nM) is continuously perfused to the recorded cell through a bath (U-tube) to get Ca^{2+} oscillation response (as a baseline). Then, the GW is added to the bath perfusion in the presence of ACh. Finally, the GW is washed out with the same concentration of ACh. With this protocol, the ACh is continuously perfused throughout the recording period, and we can compare the change of ACh-induced Ca²⁺ oscillations before GW perfusion (baseline), during GW perfusion, and after GW washout in the same recorded cell. For statistical analysis of the effects of GW on ACh-induced Ca²⁺ oscillations, we measured baseline oscillations as the charge (current area/Cm/min)¹⁸ and compared the changes of Ca²⁺ oscillations during GW perfusion and after washout of GW to the baseline. Our data showed that in the continuous presence of 10 nM ACh, 10 µM GW reduced Ca²⁺ oscillations, and this inhibitory effect was reversed after washout (Fig. 2B). A similar inhibitory effect by GW (100 μ M) was also observed on 100 nM ACh-induced Ca²⁺ oscillations using confocal Ca²⁺ imaging (Fig. 2C). Statistical analysis of the Ca²⁺ oscillation signal from 8 cells tested showed that GW significantly reduced ACh-induced Ca^{2+} oscillations from baseline level of -4.69 ± 0.32 to -1.68 ± 0.32 nC/min (the level after GW exposure, n = 8, paired *t* test *p* < 0.001, Fig. 2D). Ca²⁺ imaging experiments also showed a similar inhibition of Ca^{2+} oscillations by GW (n = 66, paired *t* test *p* < 0.001, Fig. 2E). After



Figure 1. Identification of CB₂R expression in mouse pancreatic acinar cells. (A) Western blot assay shows that a high-density CB₂-immunoreactive band is detected in both spleen and striatal tissues in WT and CB₁-KO mice, but is undetectable in striatal tissues or substantially reduced in CB₂-rich spleen tissues in CB₂-KO mice. (B) Immunohistochemical assays show high densities of CB₂-immunostaining in spleen slices of WT mice, which are undetectable or substantially diminished in CB₂-KO mice. (C) Immunocytochemical assays use mouse CB₂R antibody (NIDA-5633). The bright field photograph (Ca) shows freshly dissociated pancreatic acinar cells. CB₂-immunostaining (mCB₂-ir) in single dissociated pancreatic acinar cells illustrates the high densities of CB₂R proteins (Cb). DAPI staining demonstrates cell nucleus (Cc). The "Merged" image shows superimposed mCB2-ir and DAPI images (Cd).

washout of GW, Ca^{2+} oscillations were partially recovered in both patch recording and Ca^{2+} imaging. These results suggest that activation of CB_2R by GW inhibits ACh-induced intracellular Ca^{2+} signals in freshly isolated pancreatic acinar cells.

GW Inhibits ACh-Induced Ca²⁺Oscillations in a Concentration-Dependent Manner. To profile the pharmacological effect of GW on ACh-induced Ca²⁺ oscillations, we examined the effects of different concentrations of GW on 10 nM ACh-induced Ca²⁺ oscillations. Figure 3A–C show that GW inhibited Ca²⁺ oscillations in a concentration-dependent manner. In 1 µm GW group, Ca²⁺ oscillation levels were slightly reduced from baseline -4.59 ± 1.11 to -4.46 ± 1.25 nC/min (p > 0.05, n = 8). In 10 µm GW group, Ca²⁺ oscillation levels were reduced from baseline -4.69 ± 0.32 to -1.68 ± 0.32 nC/min (p < 0.0001, n = 8). In 100 µm GW group, Ca²⁺ oscillation levels were reduced from baseline -5.77 ± 1.75 to -0.50 ± 0.15 nC/min (p < 0.05, n = 5). Further comparisons determined that Ca²⁺ oscillation levels differed significantly between the following groups: GW 1 µM vs. $10 \mu M$ (p < 0.05), GW 1 µM vs. $100 \mu M$ (p < 0.05), and GW 10 µM vs. $100 \mu M$ (p < 0.01), which confirms that GW inhibition occurs in a concentration-dependent manner.

GW Inhibits ACh-Induced Ca²⁺ Oscillations by a Selective Action on CB₂Rs. To address the question of whether GW inhibition of ACh-induced Ca²⁺ oscillations is mediated through CB₂Rs, we designed three sets of experiments. 1) We tested the effect of a selective CB₂R antagonist (AM630) on GW inhibition of Ca²⁺ oscillations. 2) We examined GW inhibitory effects on pancreatic acinar cells prepared from CB₁-KO and CB₂-KO mice. 3) We evaluated the effects of a selective CB₁R agonist (ACEA) on ACh-induced Ca²⁺ oscillations. The results of these experiments demonstrated that GW inhibition of ACh-induced Ca²⁺ oscillations was presented in WT (Fig. 4A) and CB₁R-KO mice (Fig. 4B), but was absent in CB₂R-KO mice (Fig. 4C). Figure 4D summarizes pooled data demonstrating the effect of GW on 30 nM ACh-induced Ca²⁺ oscillations in WT (p < 0.01, n = 5), CB₁R-KO (p < 0.001, n = 6), and CB₂R-KO (p > 0.05, n = 8) mice. Furthermore, co-application of AM630 (0.1 μ M) and GW (10 μ M) abolished the inhibitory effect of GW on 10 nM ACh-induced Ca²⁺ oscillations (baseline vs. AM630 + GW p > 0.05, n = 10), while AM630 alone had no affect (baseline vs. AM630, p > 0.05, n = 10, Fig. 4E). Finally, we found that CB₁R agonist, ACEA (10 μ M) also reduced ACh-induced Ca²⁺ oscillations but this effect was likely mediated through ethanol that was used to dissolve ACEA (Supplemental Fig. 3). Together, these results suggest that GW inhibits ACh-induced intracellular Ca²⁺ signaling through the action of CB₂Rs.

GW Inhibits ACh-Induced Ca²⁺ Oscillations through Membrane CB₂Rs. Our data clearly demonstrated that GW inhibited ACh-induced intracellular Ca^{2+} oscillations. However, it remained unclear whether



Figure 2. Effects of the CB₂R agonist on ACh-induced Ca²⁺ oscillations in dissociated pancreatic acinar cells. (A) Experimental protocol shows continuous exposure to ACh (baseline), addition of GW on top of ACh, and washout of GW (with ACh). A typical trace of ACh-induced Ca²⁺ oscillations measured using patch-clamp whole-cell recording in voltage-clamp mode (measuring Ca²⁺-dependent Cl⁻ current). In the continuous presence of ACh (10 nM), addition of GW (10 μ M) reversibly reduced Ca²⁺ oscillations. (B) A typical trace of ACh-induced Ca²⁺ oscillations. (B) A typical trace of ACh-induced Ca²⁺ oscillations. (B) A typical trace of ACh-induced Ca²⁺ oscillations. Statistical analysis shows that GW significantly reduces ACh-induced Ca²⁺ oscillations in both patch-clamp recording (C) and Ca²⁺ imaging experiments (D). (D) The net charge of ACh-induced baseline Ca²⁺ oscillations (prior to GW application) is compared to the charge during GW application (+GW) and during washout of GW (Washout). Numbers in parentheses indicate the number of cells tested. Columns indicate the mean of current charge ± SEM (left) and the mean DF/DO ± SEM (right) as compared to the baseline level. ***Indicates *p* < 0.001 for the value compared to baseline level. Statistic comparison between the levels of baseline and washout of GW showed significance (*p* < 0.05) in patch-clamp data (Fig. 2D left panel) and in Ca²⁺ imaging data (*p* < 0.01, Fig. 2D right panel).

GW inhibition was mediated through extracellular or intracellular CB_2Rs . GW could act on extracellular membrane CB_2Rs and/or modulate muscarinic receptors, or GW could affect intracellular CB_2Rs , and then modulate signal molecules such as G-protein and/or inositol 1,4,5-trisphosphate (IP₃) receptors²¹. To distinguish among these possibilities, we designed two experiments, in which, either the CB_2R agonist (GW) or antagonist (AM630) was applied internally or in which, IP₃ was applied internally. When GW (100µM) was added into the recording electrode and a perforated whole-cell recording (amphotericin B) was performed, bath-application of 10 nM ACh induced Ca²⁺ oscillations. When the recording mode was switched from perforated to conventional



Figure 3. GW inhibits ACh (10 nM)-induced Ca²⁺ oscillations in a concentration-dependent manner. Typical traces show the effect of different concentrations of GW: (A) 1 μ M, (B) 10 μ M, (C) 100 μ M. (D) Bar graph summarizes the concentration-dependent effect of GW on ACh-induced Ca²⁺ oscillations. The number of cells tested is stated for each condition in parentheses. Columns show the mean of charge ± SEM. *Indicates p < 0.05, ***Indicates p < 0.001 for the values between baseline level of ACh response indicated as open columns at left and the level after GW exposure (solid columns). No asterisk mark means (GW 1 μ M group) p > 0.05.

whole-cell recording by a brief suction, GW was infused into the recorded cell, and no detectable inhibitory effect on ACh-induced Ca²⁺ oscillations was present (Fig. 5A,D). Using the same experimental protocol, we applied AM630 (1 μ M) intracellularly and found that internal AM630 failed to prevent bath-applied GW-induced inhibition in the ACh-induced Ca²⁺ oscillations (Fig. 5B,D). In the presence of intracellularly applied IP₃ (30 μ M), which causes IP₃-induced Ca²⁺ oscillations, GW produced little inhibitory effect on the IP₃-induced Ca²⁺ oscillations (Fig. 5C,D). These data suggest that GW inhibition of ACh-induced Ca²⁺ oscillations is not mediated through intracellular IP₃ receptors. Together, these results suggest that GW inhibition of intracellular Ca²⁺ oscillations is mediated through CB₂Rs on the surface of the cytoplasmic membrane.

Effects of GW on CCK-Induced Ca²⁺ Oscillations. Data presented thus far demonstrate that GW inhibited ACh-induced Ca²⁺ oscillations through cell membrane CB₂Rs, perhaps through CB₂Rs and muscarinic receptor cross talk. To test this possibility, we applied CCK to induce Ca²⁺ oscillations, which occurs through different receptor signaling pathway than muscarinic receptor, and examined the effects of GW on the CCK-induced



Figure 4. GW (10 µM) inhibits ACh-induced Ca²⁺ oscillations through CB₂Rs. (A) A typical trace shows the effect of GW on ACh (30 nM)-induced Ca²⁺ oscillations in WT mice cells. (B) GW fails to inhibit ACh-induced Ca²⁺ oscillations in acinar cells prepared from CB₁R-KO mice. (C) GW inhibits ACh-induced Ca²⁺ oscillations in acinar cells prepared from CB₂R-KO mice. (D) Columns show the mean of charge \pm SEM, summarizing the effect of GW on ACh-induced Ca²⁺ oscillations in WT, CB₁R-KO and CB₂R-KO mice cells. The number of cells tested is stated for each condition in parentheses. **Indicates *p* < 0.01 compared to the baseline level of ACh response (open columns) to the level after GW exposure (solid columns). (E) Bar graph demonstrates that the CB₂R antagonist (AM630) alone does not significantly affect ACh-induced Ca²⁺ oscillation response (baseline vs. AM630: *p* > 0.05) but abolishes GW-induced inhibition (baseline vs. Am630 + GW: *p* > 0.05).

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Ca²⁺ oscillations. As shown in Fig. 6, bath application of 10 pM CCK induced Ca²⁺ oscillation responses, which were not affected by bath application of GW (100 μ M, Fig. 6A). In the same recorded cell, bath application of GW (100 μ M) dramatically inhibited 10 nM ACh-induced Ca²⁺ oscillations (Fig. 6B). Figure 6C summarizes pooled data from 4 cells tested, and no significant effect of GW on CCK-induced Ca²⁺ oscillations was found (p > 0.05, n = 4, Ca), but GW inhibited ACh-induced Ca²⁺ oscillations in the same recorded cell (p < 0.01, n = 4, Cb).

L-arginine Potentiates ACh-Induced Ca²⁺ Oscillations. L-arginine is used to induce acute pancreatitis in rodents²². In dissociated pancreatic acinar cells, bath-application of L-arginine for 10 min enhanced ACh-induced Ca²⁺ oscillations from baseline level of 4.93 ± 0.39 to 10.34 ± 1.83 nC/min (Fig. 7Aa,b), which was not reversible after washout for 10 min (Ca²⁺ oscillations between L-arginine exposure and washout groups p > 0.05, n = 6, Fig. 7Ac). Statistical analysis revealed that L-arginine significantly enhanced ACh-induced Ca²⁺ oscillations (p < 0.05) in an irreversible manner (Fig. 7B).



Figure 5. GW inhibits **ACh-induced Ca**²⁺ oscillations through cytoplasmic, rather than intracellular, **CB**₂**Rs.** (**A**) A typical trace shows ACh-induced Ca²⁺ oscillations between perforated and conventional whole-cell recordings, in which the pipette solution contained 100 μ M GW. Infusion of GW into the recorded cell does not reduce ACh-induced Ca²⁺ oscillations (**Aa**). Ab: Summary of pooled data, demonstrating no significant difference between ACh responses with and without intracellular GW (p > 0.05, n = 6). (**B**) Internal infusion of AM630 fails to prevent bath-applied GW inhibition of ACh-induced Ca²⁺ oscillations. **Indicates p < 0.01. (**C**) Intracellular applied IP₃-induced Ca²⁺ oscillations are not sensitive to bath-applied GW (Ca). (Cb) Bar graph summarizes the effect of GW on IP₃-induced Ca²⁺ oscillations and showing no significance before and after GW exposure (p > 0.05, n = 7). The number of cells tested is stated for each condition in parentheses.

GW Prevents L-arginine-Enhanced Ca²⁺ Oscillations. Next, we sought to determine whether GW could eliminate L-arginine-induced enhancement of Ca^{2+} oscillations. We showed that either pre-treatment with GW (Fig. 8A), or co-administration of GW (10 μ M) and L-arginine (Fig. 8B), abolished L-arginine-induced enhancement of Ca^{2+} oscillations (Fig. 8C,D), suggesting that selective activation of acinar cell CB₂Rs significantly eliminates L-arginine-induced enhancement of intracellular Ca²⁺ signals in mouse pancreatic acinar cells.

GW Improves L-arginine-Induced Pathology. Finally, we tested whether systemic injection of GW can prevent L-arginine-induced elevation of Ca^{2+} oscillations, and subsequent pathological changes including enhancement of pancreatic amylases (AMS) and pulmonary peritoneal macrophages (MPO) levels, which are two major effects present in early-stage of acute pancreatitis. We injected L-arginine (4.0 g/kg, i.p.) to establish an acute pancreatitis model^{23,24}, and dissociated pancreatic acinar cells 24 hours later, then compared ACh-induced Ca²⁺ oscillations between saline- and L-arginine-treated groups using Ca²⁺ imaging. Systemic L-arginine injection enhanced ACh-induced Ca^{2+} oscillations compared to systemic saline injection, but GW and L-arginine co-injected showed similar level of ACh-induced Ca²⁺ oscillations (Fig. 9A). Compared to the ACh-induced Ca²⁺ oscillations in saline-treated mice, the acinar cells prepared from L-arginine-treated mice showed a significant increase in Ca²⁺ oscillation response (saline vs. L-arginine group, p < 0.01), while co-injection of GW and L-arginine reduced L-arginine's effect (saline vs. L-arginine + GW group, p > 0.05). These results suggest that the activation of pancreatic acinar cell CB₂Rs may prevent early pathogenesis of acute pancreatitis through the inhibition of enhanced intracellular Ca²⁺ signals. In addition, co-injection of GW (10 mg/kg, i.p.) and L-arginine (4g/kg, i.p.) also significantly reduced pancreatic L-arginine-induced enhancement of AMS (saline vs. L-arginine, p < 0.05, and saline vs. GW + L-arginine, p > 0.05; Fig. 9C) and pulmonary MPO levels (saline vs. L-arginine, p < 0.05, and saline vs. GW + L-arginine, p > 0.05; Fig. 9D). These results suggest that the activation of pancreatic acinar cell CB₂Rs may prevent early pathogenesis of acute pancreatitis through the inhibition of intracellular Ca²⁺ signals.

Discussion

The novel findings of this study are that the activation of membrane CB_2Rs by GW reduces ACh-, but not CCK-induced intracellular Ca^{2+} oscillations, and GW induced reduction of Ca^{2+} oscillations in a



Figure 6. Effects of GW on CCK-induced Ca²⁺ oscillation. (A) Representative typical trace of CCK (10 pM)induced Ca²⁺ oscillations, which are not affected by 100 μ M GW. (B) In the same recorded cell, ACh (10 nM)induced Ca²⁺ oscillations are completed eliminated by GW. (C) Bar graph summarizes the effect of 100 μ M GW on CCK (Ca)- and ACh (Cb)-induced Ca²⁺ oscillations. No asterisk mark (CCK vs CCK + GW 100 μ M) means p > 0.05 (n = 4). **Indicates p < 0.01 compared between ACh 10 nM and ACh + GW 100 μ M, n = 4.

concentration-dependent manner. The CB₂R-mediated reduction of ACh-induced Ca²⁺ oscillations is abolished by pharmacological blockade of CB₂Rs (AM630) or is absent in CB₂-KO mice, but not in CB₁-KO mice. The pancreatitis inducer, L-arginine, significantly enhances ACh-induced intracellular Ca²⁺ oscillations, and the CB₂R agonist, GW, abolishes this L-arginine effect. In addition, this CB₂R agonist also improved L-arginine-induced pathological changes. Collectively, our data demonstrate that CB₂R agonist GW reduces ACh-enhanced intracellular Ca²⁺ signals in mouse pancreatic acinar cells, and this may underlie an important cellular mechanism for a CB₂R agonist to serve as a new candidate for treating acute pancreatitis.

CB₂**R Expression in Mouse Pancreatic Acinar Cells.** Previously, in rodent pancreatic acinar cells, CB_2R protein expression was found using immunohistochemical staining and Western blot^{10,25}. In mouse pancreatic tissue, both CB_1R and CB_2R mRNA were identified using real-time RT-PCR and immunohistochemical staining¹⁰. In the present study, we confirmed that CB_2R proteins were expressed in freshly isolated mouse pancreatic acinar cells, which is consistent with previous report¹⁰. Our data demonstrate that CB_2R are expressed in mouse pancreatic acinar cells and they may play an important role in modulating acinar cells function.

CB₂**R Agonist Reduces ACh-Induced Ca**²⁺ **Oscillations in Mouse Pancreatic Acinar Cells.** Mouse pancreatic acinar cells have been used as an excellent cell model of agonist-induced Ca²⁺ oscillations for studying pancreatitis²⁶. We examined whether a selective CB₂R agonist, GW, affected ACh-induced Ca²⁺ oscillations in the isolated pancreatic acinar cells through CB₂Rs. Using both patch-clamp recording and confocal Ca²⁺ imaging techniques, we found that GW significantly reduced ACh-induced Ca²⁺ oscillations, and this inhibition is GW-concentration dependent. We also tested another selective CB₂R agonist, JWH-133, on the ACh-induced Ca²⁺ oscillations and the another selective CB₂R agonist, JWH-133, on the ACh-induced Ca²⁺ oscillations and the ACh-induced Ca²⁺ oscillations and the concentration dependent.



Figure 7. Effects of L-arginine on ACh-induced Ca²⁺ oscillations. (A) Representative traces of ACh-induced Ca²⁺ oscillations before (A), during (B), and after (C) bath-application of L-arginine (10 mM, L-Arg). Traces (A–C) were recorded from the same cell. (B) Bar graph summarizes the charge (±SEM) and shows an enhanced effect of L-Arg on ACh-induced Ca²⁺ oscillations. Six cells were assessed before and after L-Arg application. *Indicates p < 0.05 compared to baseline level. There was no significance between L-Arg application and washout of L-Arg (p > 0.05), suggesting the effects of L-Arg is non-reversible.

 Ca^{2+} oscillations, and found a similar inhibition (Supplemental Fig. 4), but the inhibitory effect of JWH-133 was weaker (a higher concentration of JWH-133 was needed compared with GW to induce the same inhibition). It was reported that GW acts as a potent and selective partial agonist for CB_2R with an EC_{50} of 0.65 nM and selectivity of around $1200 \times$ for CB_2R over $CB_1R^{27,28}$, while JWH-133 has an EC_{50} of 3.4 nM and selectivity of around $200 \times$ for CB_2R over CB_1R^{29} . These findings may explain why GW is more potent than JWH-133 for ACh-induced Ca^{2+} oscillations.

Accumulating evidence demonstrates a complex relationship between the cannabinoid ligand (and receptors) and intracellular Ca^{2+} signals in different types of cells. For example, on one hand, activation of cannabinoid CB₁R or CB₂R increased (initiated) intracellular Ca^{2+} levels in endothelia cells³⁰, submandibular acinar cells³¹, canine kidney cells³², and bladder cancer cells³³. On the other hand, in pancreatic beta cells, the activation of either CB₁R³⁴ or CB₂R³⁵ reduced glucose-induced intracellular Ca²⁺ oscillations and insulin release. It has been reported that anandamide reduced intracellular Ca²⁺ concentration through the suppression of a Na⁺/Ca²⁺ exchanger current in rat cardiac myocytes³⁶. To our knowledge, ours is the first report that a selective CB₂R agonist reduces intracellular Ca²⁺ signals in mouse pancreatic acinar cells. Considering that Ca²⁺ plays an important role in cellular function, especially enzyme secretion in pancreatic acinar cells, our data suggest that CB₂R modulates an important aspect of pancreatic acinar cell physiology and pathophysiology.

 CB_2R Agonist Reduces ACh-Induced Ca^{2+} Oscillations through Membrane CB_2Rs . Cannabinoid ligands exert their pharmacological effects through CB_1R or CB_2R , but in some cases they also can act on non-cannabinoid targets³⁷. We determined whether GW modulated intracellular Ca^{2+} signals through a cell membrane or cytosolic CB_2Rs . First, we examined the effects of pharmacological manipulations of CB_1R and



Figure 8. Effects of GW on L-Arg induced enhancement of Ca^{2+} oscillations. (A) After pretreatment with GW, bath-applied L-Arg (10 mM for 10 min) fails to enhance ACh-induced Ca^{2+} oscillations. Traces in Fig. 8Aa,b were recorded from the same cell. (B) Without pretreatment, bath-applied L-Arg enhances ACh-induced Ca^{2+} oscillations, and under this condition, the addition of GW also reduces L-Arg-induced enhancement of Ca^{2+} oscillations. Traces in Fig. 8Ba,b were recorded from the same cell. (C,D) GW significantly blocks L-Arg-induced enhancement of Ca^{2+} oscillations either with or without pretreatment of GW. Bar graphs represent averaged charge \pm SEM. The number of cells tested is stated for each condition in parentheses. *Indicates p < 0.05, ***Indicates p < 0.001.

 CB_2R and found that the CB_2R selective antagonist AM630 abolished GW-induced reduction of Ca^{2+} oscillations, suggesting that GW modulates ACh-induced Ca^{2+} oscillations through the CB_2Rs . Then, we genetically manipulated cannabinoid receptors and compared the effects of GW on Ca^{2+} oscillations between WT and CB_2R -KO mice, and also WT and CB_1R -KO mice. We found that in CB_2R -KO but not CB_1R -KO mice, GW lost its inhibitory effect, further confirming that CB_2R is the key target for mediating GW-induced reduction in Ca^{2+} oscillations.

In a group of cells tested, we found that a CB_1R agonist, ACEA (dissolved by ethanol; 10- μ M ACEA solution contained 7.3-mM ethanol) reduced ACh-induced Ca²⁺ oscillations (Supplemental Fig. 3); however, the control experiments using the same concentration of ethanol (7.3 mM) also reduced ACh-induced Ca²⁺ oscillations, and the inhibitory effect of ACEA was not absent in the acinar cells dissociated from CB₁R-KO mice, suggesting a non-specific effect, likely caused by ethanol. In addition, we also tested the effects of DMSO (GW was dissolved by DMSO to 100 mM stock solution), and found that 1 μ M DMSO itself did not affect ACh-induced Ca²⁺ oscillations (Supplemental Fig. 5). Together, our data support the conclusion that GW selectively acts on acinar cell CB₂Rs and reduces ACh-induced Ca²⁺ oscillations.





Finally, we asked where the CB₂Rs are located (membrane or cytosolic CB₂Rs). To address this question, we designed three experiments. We first examined the effect of bath-applied GW on the Ca²⁺ oscillations induced by intracellular application of IP₃, and found that GW did not affect IP₃-induced Ca²⁺ oscillations, suggesting that the target that mediated GW-induced inhibition in Ca²⁺ oscillations is located in the signal pathway before IP₃ receptors, and not on the IP₃ receptor itself. We then intracellularly applied GW through a recording electrode to examine the effect of intracellular administration of GW on bath ACh-induced Ca²⁺ oscillations. Finally, we intracellular infusion of GW (even at 100 µM) did not alter ACh-induced Ca²⁺ oscillations. Finally, we intracellularly applied AM630 through a recording electrode to examine the effect of bath-applied GW on ACh-induced Ca²⁺ oscillations. Our data showed that intracellular infusion of AM630 did not prevent bath-applied GW-induced reduction of Ca²⁺ oscillations. Collectively, our data support the conclusion that GW modulates intracellular Ca²⁺ signaling through the membrane CB₂Rs in pancreatic acinar cells.

Possible Mechanisms of GW-Induced Reduction in ACh-Induced Ca²⁺ Oscillations. The precise mechanism by which GW modulates intracellular Ca²⁺ signals is unclear. Our data show that membrane CB₂Rs are necessary for mediating GW's effect. GW's action in ACh-induced Ca²⁺ oscillations should occur at the G-protein-mediated signal pathway between muscarinic receptor (M₃) activation and IP₃ production because GW did not affect IP₃-induced Ca²⁺ oscillations. We also demonstrated that GW failed to affect ACh-induced Ca²⁺ oscillations in pancreatic acinar cells prepared from CB₂R-KO mice, suggesting that GW likely did not affect muscarinic receptor function. In addition, we found that bath-applied GW failed to inhibit CCK-induced Ca²⁺ oscillations even at 100 µM, suggesting that GW selectively modulates muscarinic receptor-mediated G-protein signaling³⁸. Therefore, the possible mechanisms for GW-induced modulation of ACh-induced Ca²⁺ oscillations may involve cross talk between muscarinic receptor- and CB₂R-mediated G-protein signal pathways, such as homologous and/or heterologous desensitization of GPCRs, the activation of one type of GPCR can rapidly terminate another GPCR signaling through the internalization of receptors after binding, phosphorylation of G-protein coupled receptor kinases, and formation of complexes with β-arresting^{39,40}. In addition, the activation of a GPCR may also result in temporary inhibition of another GPCR signal through a heterologous desensitization, which does not involve receptor internalization, but activation of several signal transduction pathways, particularly protein kinase C (PKC)- and PKA-dependent signaling pathways^{38,41}. It has been reported that intracellular cyclic AMP-generated substances play an important role in regulation of IP₃ and Ca²⁺ responses to ACh in rat submandibular acini. Investigators found that intracellular cAMP increased IP₃ formation in response to ACh, while blocking PKA by H89 reduced IP₃ formation⁴¹. Because it is well known that the activation of CB₂Rs significantly reduces intracellular cAMP levels, we thus postulated that GW may activate CB₂Rs, reduce cAMP, and in turn reduce intracellular IP₃ production, and lead to a reduction of ACh-induced Ca²⁺ oscillations. Our findings warrant further testing of this hypothesis.

Clinical Significance of CB₂R-Mediated Reduction of Ca²⁺ Oscillations in Pancreatic Acinar

Cells. Pancreatic acinar cells are functional units of the exocrine pancreas. They synthesize, store, and secrete inactive preforms of digestive enzymes into the lumen of the acinus. The activity of pancreatic acinar cells is crucially modulated by the secretagogues ACh and CCK; both can act on their specific membrane receptors (muscarinic and CCK receptor, respectively) and then induce an elevation in cytoplasmic calcium. If high concentrations of intracellular Ca²⁺ persist, intracellular signaling is disrupted, cell damage occurs, and acute pancreatitis forms. Emerging evidence suggests that the earliest abnormalities of acute pancreatitis arise by aberrant elevation of intracellular Ca^{2+} within acinar cells because the sustained intracellular Ca^{2+} elevation activates intracellular digestive proenzymes resulting in necrosis and inflammation, and pharmacological blockade of store-operated or Ca^{2+} release-activated Ca^{2+} channels would prevent sustained elevation of intracellular Ca^{2+} , and consequence protease activation and necrosis³. In the present study, we provide the first evidence that the CB₂R agonist, GW, reduces ACh-induced Ca²⁺ oscillations, abolishes L-arginine–induced enhancement of Ca²⁺ oscillations and prevents L-arginine-induced elevation of both pancreatic AMS and pulmonary MPO levels. These results suggest that a CB₂R agonist may serve as a novel therapeutic strategy to prevent and/or treat acute pancreatitis. This conclusion is consistent with previous report that a CB₂R agonist exhibits a protective effect on pathogenesis in an acute pancreatitis animal model¹⁰. Our data showing a reduction of intracellular Ca²⁺ signaling by GW also provide a new target to interpret the role of CB₃R agonists in treating acute pancreatitis in addition to CB₂R-mediated anti-inflammation.

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Author Contributions

All authors reviewed the manuscript. Z.H. reached the literature, designed patch-clamp experiments, performed most patch-clamp experiments, collected patch-clamp data, analyzed patch-clamp data, and prepared Figures 2-5, 7 and 8. HW designed the acute pancreatitis experiments, collected and analyzed data, prepared Figure 9, and wrote part of the manuscript. M.Z. researched the literature, performed Ca^{2+} imaging experiments, collected and analyzed data, prepared Figures 2B,D and 9A,B. NS researched the literature, performed acute pancreatitis experiments, collected and analyzed data, and prepared Figure 9. F.S. researched the literature, performed acute pancreatitis experiments, collected and analyzed data, and prepared Figure 9. J.S. researched the literature, designed Ca²⁺ imaging experiments, analyzed data, prepared Figures 2B,D and 9A,B and wrote part of the manuscript. H.Z. designed and performed immunocytochemical experiments, analyzed data, prepared Figure 1, and wrote part of the manuscript. Z.X. designed molecular and cell biological experiments, analyzed data, prepared Figure 1 and revised the manuscript. Q.L. designed molecular biological experiments, analyzed data, prepared Figure 1 and revised the manuscript. K.X. and D.C. performed patch-clamp experiments and collected data for Figure 6. M.G. researched the literature, performed some patch-clamp experiments, collected patch-clamp data, analyzed patch-clamp data, and participated in the preparation of Figures 2-5. R.P.H. designed and advised on patch-clamp experiments and revised the manuscript. X.F. participated in the design of all experiments, discussed and analyzed data, wrote part of the manuscript, and revised the manuscript. J.W. researched the literature, designed experiments, analyzed data, finalized all figures, and wrote the main manuscript text.

Additional Information

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