The Feverfew plant-derived compound, parthenolide enhances platelet production and attenuates platelet activation through NF-κB inhibition

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A B S T R A C T
Introduction: Few treatments are available that can safely and effectively stimulate new platelet production for thrombocytopenic patients. Additionally, recipients of transfused platelets may experience an inflammatory response due to stored platelets becoming unnecessarily activated, thus creating the need for suitable agents that will dampen undesirable platelet activation. We investigated the effect of the feverfew plant-derived compound, parthenolide on platelet production and platelet activation because of its well-studied ability to induce apoptosis or differentiation in some types of cancer.

Methods: Parthenolide was used to treat human megakaryoblastic cell lines, primary human and mouse megakaryocytes. Resulting platelet production and function was measured via flow cytometry. The two most common parthenolide signaling mechanisms, oxidative stress and nuclear factor-κB inhibition, were assessed within the megakaryocytes using reactive oxygen species, glutathione and luciferase reporter assays. The influence of parthenolide on ex vivo platelet activation was tested with parthenolide pretreatment followed by collagen or thrombin activation. The resulting P-selectin surface expression and released soluble CD40 ligand was measured.

Results: Parthenolide stimulates functional platelet production from human megakaryocyte cell lines, and from primary mouse and human megakaryocytes in vitro. Parthenolide enhances platelet production via inhibition of nuclear factor-κB signaling in megakaryocytes and is independent of the parthenolide-induced oxidative stress response. Additionally, parthenolide treatment of human peripheral blood platelets attenuated activation of stimulated platelets.

Conclusion: Overall, these data reveal that parthenolide has strong potential as a candidate to enhance platelet production and to dampen undesirable platelet activation.

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Introduction
Platelets play a fundamental role in hemostasis and have emerged as important contributors to inflammation and immunity [1–3]. Thrombocytopenia, a disorder of low platelet counts and sometimes abnormal bleeding, is a serious problem that has limited treatment options, the most widely used being platelet transfusion. There is great interest in discovery of new compounds to boost thrombopoiesis. We previously reported that 15-deoxy-Δ12,14-Prostaglandin J2 is capable of enhancing platelet production via oxidative stress [4]. In screening other thrombopoiesis-enhancing candidates, the anti-inflammatory feverfew plant-derived compound, parthenolide showed strong potential. Parthenolide is currently being investigated for its ability to induce apoptosis [5–8] or differentiation [9] in some types of cancer cells. The best known mechanisms of parthenolide are the inhibition of nuclear factor-κB (NF-κB) activity via inhibitor-κB kinase [10] and the induction of oxidative stress [5,11]. Interestingly, both the increase of reactive oxygen species (ROS) and decrease of NF-κB signaling have been correlated with the final stages of megakaryocyte maturation and thrombopoiesis [12,13]. We therefore chose to further investigate the mechanism of parthenolide enhancement of platelet production to better understand how parthenolide and other drug candidates could serve as thrombopoietic agents.

There are 150–400 × 10^6 platelets per μL of whole human blood, and these cells are important early responders to stress signals, playing an intricate role in the initiation and maintenance of inflammation. Activated platelets upregulate surface immunomodulatory proteins such as P-selectin and CD40 ligand (CD40L) [14] and
release many inflammatory mediators from their α-granules and dense bodies such as platelet factor-4, prostaglandin-E$_2$, serotonin, vascular endothelial growth factor, RANTES, and soluble CD40 ligand (sCD40L) [1, 2, 15]. These mediators can act to amplify an inflammatory or immune response, and contribute to diseases such as type-2 diabetes. Overabundance of inflammation can be detrimental to the vasculature or tissue where it is occurring. Thus, there is a rising demand for anti-platelet molecules that would decrease unwanted vasculature or tissue where it is occurring. Thus, there is a rising demand for anti-platelet molecules that would decrease unwanted vasculature or tissue where it is occurring. Parthenolide has previously been reported to have some anti-platelet activity [16–18]. However, the exact mechanism of platelet inhibition by parthenolide has not been elucidated. We and others have recently demonstrated that functional NF-κB family proteins are not only present in megakaryocytes and platelets, but play a role in their activation [19–21]. We therefore tested whether parthenolide would influence platelet activation via NF-κB.

Materials and methods

Reagents and antibodies

Parthenolide and the mouse anti-human heme oxygenase-1 (HO-1) antibody were purchased from Biomol (Plymouth Meeting, PA); BMS-345541, Tyrodes Salt Solution, fibroinogen and thrombin from Sigma (St. Louis, MO); Carboxy-H$_2$DCFDA from Invitrogen (Carlsbad, CA); Collagen from Chrono-log Corporation (Havertown, PA); Cignal Lenti NfκB Reporter (luc) Kit: CLS-013 L from Qiagen (Raleigh, NC); Alexa Fluor 488 anti-human CD61:Alexa Fluor 647 from AbD Serotec (Raleigh, NC); Alexa Fluor 488 anti-human CD62P from Biolegend (San Diego, CA); BIT 9500 from StemCell Technologies (Vancouver, Canada); Recombinant Human: IL-1β, IL-6, thrombopoietin (TPO) and Stem Cell Factor (SCF) were purchased from R&D Systems (Minneapolis, MN); mouse anti-actin from Oncogene Research (Cambridge, MA).

Cell line culture conditions

Meg-01 and MØ7e cells were purchased from the American Type Culture Collection (Rockville, MD), and cultured in RPMI 1640 (Invitrogen) with 5% fetal bovine serum (Invitrogen), 10 mM HEPES (Sigma), 2 mM L-glutamine (Invitrogen), 4.5 g/L glucose (Invitrogen), and 50 μg/mL gentamicin (Invitrogen). MØ7e cells were additionally supplemented with 100 ng/mL of granulocyte macrophage colony-stimulating factor (R&D Systems). All treatments were diluted to a 10X concentration in Phosphate Buffered Saline (Invitrogen) before added to culture. Vehicle treatments contain equal volume of DMSO to the maximum concentration of drug treatments. NF-κB luciferase reporter cell lines were obtained by transducing 1x 10⁵ Meg-01 or MØ7e cells with 1x 10⁶ transducing units for 5 hours. Positively transduced cells were selected for and continuously maintained in media supplemented with 2 μg/mL Puromycin (Invitrogen).

Primary Megakaryocyte differentiation and culture

Bone marrow was isolated from C57BL/6 mice (Jackson Laboratories, Bar Harbor, ME). 1x 10⁶ cells/mL were cultured in IMDM (Invitrogen) with 20% BIT 9500 and 100 ng/mL of rhTPO for 5 days, after which the cells were washed and treated with 10 μM parthenolide. Adapted from previous procedures [4], human CD34+/cord blood cells were purchased from Allcells (Emeryville, CA) and first plated at 2.5 x 10⁴ cells/mL in IMDM supplemented with 20% BIT 9500, 0.8% Low Density Lipoprotein (L-8292, Sigma), 100 ng/mL rhTPO, 50 ng/mL rhSCF, 10 ng/mL rhIL-6 and 10 ng/mL IL-1β. After 7 days of expansion, cells were cultured in the same conditions minus rhSCF, and on day 14 were used for treatment.

Megakaryocyte and platelet imaging, counting and analysis

Live megakaryocytes in culture were examined by phase-contrast microscopy with an inverted microscope (Olympus IX81), and images were captured and edited with SPOT RT Software 4.6 (New Hyde Park, NY). Cultures were stained with an anti-CD61 antibody and analyzed on an Accuri C6 flow cytometer (Ann Arbor, MI) for platelet counting and functional assays. Files were analyzed using FlowJo software (Treestar, Ashland, OR). Platelets were characterized by size gating, CD61 expression, and absence of DNA staining. Non-spun, unaltered culture was collected and stained; platelet counts were normalized to volume of analysis.

Human blood platelet isolation, treatment and analysis

Human blood was obtained from consenting donors in accordance with the Declaration of Helsinki under the University of Rochester IRB approval via venipuncture into tubes containing 3.2% sodium citrate solution (369714, BD Biosciences, Franklin Lakes, NJ). Tubes were centrifuged for 15 minutes at 250xg. Platelet rich plasma was removed centrifuged for 10 minutes at 1000xg. Platelets were washed with Tyrode’s Salt Solution: acid-citrate-dextrose (25:3) supplemented with 1 μg/mL PGE$_2$ (Cayman Chemical, Ann Arbor, Michigan), and spun again for 10 minutes, 1000xg. Platelets were resuspended in Tyrode’s Salt Solution, counted, and used immediately for treatments at a concentration of 1-3 x 10⁹ platelets/mL. Treated and activated platelets were pelleted so the supernatant could be analyzed for CD40L via ELISA (R&D Systems). Platelets were then stained with CD61 Alexa Fluor 647 and CD62P Alexa Fluor 488 antibodies and analyzed on an Accuri C6. Alternatively, platelets were treated and spread for 45 minutes onto coverslips precoated overnight in 100 μg/mL of fibronogen and blocked for 1 hour with 0.5% gelatin:BSA (Sigma A-8806). Coverslips were then washed with PBS and mounted onto slides (VWR) with Fluoromount-G (Southern Biotech, Birmingham, AL). Slides were analyzed with the Olympus BX51 light microscope and images were captured and analyzed with SPOT RT software.

Western Blotting

Whole cell lysates were prepared using ELB buffer (15 mM HEPES (pH 7)), 250 mM NaCl, 0.1% Nonidet P-40, 5 mM EDTA, 10 mM NaF, 0.1 mM Na$_2$VO$_4$, 50 μM ZnCl$_2$, supplemented with 0.1 mM PMSF, 1 mM DTT and a mixture of protease and phosphatase inhibitors. Total protein was quantified with bichinchoninic acid protein assay kit (Pierce, Rockford, IL). 25 μg of total protein per treatment was loaded and run on 12% SDS PAGE gels, electroblotted onto a Immoblot PVDF membrane (BioRad, Hercules, CA). Blots were blocked with 10% nonfat dry milk in 0.1% Tween 20 for 1 hour RT. In 2.5% milk, 1:1000 HO-1 or 1:10,000 actin primary antibodies, followed by washing and an appropriate type and amount of secondary antibodies. Blots were visualized using Immobilon Western chemiluminescence substrate (Millipore, Billerica, MA) and developed on Classic x-ray film (Laboratory Product Sales, Rochester, NY).

NF-κB TransAM assay

The TransAM assay kit was purchased from Active Motif (Carlsbad, CA), and assays were performed as indicated by the manufacturer’s protocol. Experiments were performed with duplicate samples.

Reactive oxygen species production

Meg-01 cells were treated in the dark with 10 μM carboxy-H$_2$DCFDA for 20 minutes, 37 °C, washed with PBS, and distributed for further treatment in the dark for up to 3 hours and immediately analyzed on an Accuri C6 flow cytometer.
Measurement of Intracellular GSH

Meg-01 cells were treated with 10 μM parthenolide or BSO for 0.5, 1, 3, or 6 hours. Measurements of intracellular GSH were performed as previously described [22].

Data Representation

Experimental results represent the mean +/- standard error bars. Graphs are representative of triplicates within one experiment, and all experiments were repeated independently at least 3 times. Statistical analyses are as described in figures and text.

Results

While screening multiple compounds during our investigation of thrombopoiesis-enhancing molecules, we found that the naturally occurring feverfew-derived compound, parthenolide, was capable of enhancing platelet production from the megakaryoblastic cell lines, Meg-01 and MO7e, within 24 hours (Fig. 1). The morphology of a megakaryocyte undergoing thrombopoiesis involves changes in the plasma membrane, including formation of proplatelet extensions [23]. Changes in megakaryocyte morphology towards a typical platelet-producing cell are shown in Fig. 1A and B, following parthenolide treatment in the Meg-01 and MO7e cells. The platelet-like particles produced from these cells share similar properties with freshly isolated primary platelets with expected morphology, surface markers such as CD61, and function [23].

We next determined whether this same enhancement of platelet production would occur in primary megakaryocytes after parthenolide treatment. Primary mouse megakaryocytes (PMM) were differentiated from bone marrow and treated with parthenolide. Compared to vehicle-treated PMM, parthenolide-treated PMM cultures had more megakaryocytes exhibiting a platelet-producing phenotype, such as proplatelet extensions (Fig. 2A), and there were significantly more platelets in the 24 hour treated cells (Fig. 2B). Similarly, primary human megakaryocytes (PHM) derived in vitro from human umbilical cord blood showed increased platelet-producing morphology after

![Figure 1: Parthenolide enhances platelet production from megakaryoblastic cell lines. Megakaryocytes were treated with vehicle (Veh) or 10 μM PTL. (A) Meg-01 (5 h) and (B) MO7e (3 h) cells were imaged using an inverted microscope. PTL increased the formation of proplatelet extensions compared to vehicle-treated. Black arrows indicate proplatelet extensions, which are increased in PTL-treated cells. Graphs represent mean counts of platelet-like particles per μL of (C) Meg-01 and (D) MO7e cultures at 24 hours post PTL treatment measured by the Accuri C6 Flow Cytometer. (*** indicates p < 0.001 in a post-test of one-way ANOVA).}
5 hours of parthenolide treatment, (Fig. 2C) and more platelets were produced in 24 hours compared to vehicle treated cells (Fig. 2D).

In order to characterize the mechanism for the enhanced platelet production seen after parthenolide treatment, we investigated two major pathways reported to be influenced by parthenolide: oxidative stress and NF-κB inhibition. Neither of these pathways has been studied during parthenolide treatment of megakaryocytes. Interestingly, megakaryocytes indeed undergo oxidative stress after exposure to parthenolide. Meg-01 and PHM cells increase cellular hemeoxygenase-1 (HO-1) levels in a dose and/or time-dependent fashion (Fig. 3A). HO-1 combats oxidative stress [24], and its levels have previously been shown to increase in megakaryocytes when treated with an oxidative stress inducer, 15-deoxy-Δ12,14-Prostaglandin J2 [25]. Glutathione (GSH) is another molecule critical to the redox state of the cell [26]. Total GSH levels are also increased in a time dependent manner through 6 hours of parthenolide treatment (Fig. 3B), and moreover, sustained elevation of total GSH occurs in parthenolide-treated cells through 48 hours (data not shown). BSO served as a GSH depletory control. Despite the noticeable increase in HO-1 and GSH levels, no significant reactive oxygen species (ROS) were detected in megakaryocytes at 3 hours (Fig. 3C) or any timepoint tested (data not shown). Hydrogen peroxide (H2O2) was used as a positive control to induce oxidative stress and increased the levels of ROS in Meg-01 cells (Fig. 3C). This concentration of H2O2 was not toxic to the cells (data not shown). H2O2 does not increase platelet production from Meg-01, MO7e, or PHM after 24 hours of treatment (Fig. 4). Therefore, parthenolide’s enhancement of platelet production does not appear to be caused by an oxidative stress response.

Parthenolide can inhibit NF-κB function [10]. Whether it does so in megakaryocytes is unknown. We first assessed inhibition of NF-κB using the TRANS-AM assay which quantifies the presence of specific nuclear proteins that bind to NF-κB target DNA sequences. Parthenolide treatment led to a significant decrease in the levels of p65, p50, and RelB within the nucleus of Meg-01 cells (Fig. 5A) or p65 within

Fig. 2. Parthenolide enhances platelet production from primary human and mouse megakaryocytes treated in vitro. Images of live megakaryocyte cultures taken from an inverted microscope of (A) primary mouse megakaryocytes (PMM), or (B) primary human megakaryocytes (PHM), after 2 or 5 hours, respectively, in the presence of vehicle (Veh) or 10 μM PTL. Black arrows indicate proplatelet extensions, which are increased in PTL-treated cells. Graphs show mean CD61 positive platelet-sized events per μL of (C) PMM or (D) PHM culture at 24 hours post PTL treatment, measured by an Accuri C6 Flow Cytometer. (* indicates p<0.01 in a two-tailed Student T test).
MO7e cells (Fig. 5B) by 24 hours. To support these findings, an NF-κB luciferase reporter was transduced into Meg-01 and MO7e cells. Parthenolide significantly reduced the basal NF-κB activity at 6 hours. However at 24 hours, luciferase levels returned to normal (Fig. 5C-D). A small molecule NF-κB inhibitor, BMS-345542 (BMS), was used as a positive control in the luciferase assay. We also tested BMS on the two megakaryocyte cell lines to determine if this too, enhanced platelet production. BMS dose-dependently increased platelet production from both cell lines after a 24 hour treatment (Fig. 5E-F).

Increases of platelet numbers are only clinically relevant if the platelets produced are functional. Therefore, we determined if platelets created from parthenolide-treated PHM were capable of becoming activated. Upon collagen treatment, normal human platelets demonstrate a decrease in size and increase in granularity, indicative of activation because they are releasing granules and changing morphology [27]. We isolated platelets from parthenolide-treated PHM and tested whether they were still able to be activated by collagen. Platelets formed from either vehicle- or parthenolide-treated PHM showed a similar ability to be activated by collagen, supporting their functionality (Fig. 6).

We addressed the effects of parthenolide on freshly isolated human platelets because it was previously reported that parthenolide

![Graph A](image1.png)

![Graph B](image2.png)

![Graph C](image3.png)

Fig. 3. Parthenolide generates an oxidative stress response in megakaryocytes. (A) Western blots of whole cell lysates from 6 and 24 hour PTL or 24 hour vehicle (Veh)-treated Meg-01 cells (left) or PHM (right) for hemeoxygenase-1 (HO-1) with actin as a loading control. (B) Meg-01 cells were exposed to vehicle, a known GSH depletor, buthionine sulfoximine (BSO), or PTL for the indicated times. Mean micrograms of total glutathione levels (GSH) normalized to total protein from each treatment are shown in the graph. PTL-treatment over time showed an increase of total cellular glutathione beginning at 3 hours (p<0.01 from a post-test of one-way ANOVA). (C) Meg-01 cells were prestained with H2DCFDA, which detects ROS, or unstained (US) and then treated for three hours with vehicle (Veh), 10 μM PTL, or a positive control ROS inducer, 50 μM H2O2. Cells were immediately run on an Accuri C6 flow cytometer and histograms represent H2DCFDA fluorescence. (The vehicle and PTL curves may be hard to distinguish as they exactly overlap.)

Fig. 4. The oxidative stress response from Parthenolide does not contribute to its enhancement of platelet production. Graphs illustrate mean counts of platelet-like particles per μL of (A) Meg-01, (B) MO7e, or (C) PHM culture at 24 hours post treatment of vehicle (Veh), 10 μM PTL, or 50 μM H2O2, measured by an Accuri C6 Flow Cytometer. (** indicates p<0.001; * indicates p<0.05 from a post-test of a one-way ANOVA).
decreases serotonin release from activated platelets [16], therefore showing potential as an anti-platelet therapy. Herein, we tested whether parthenolide could decrease platelet activation. Upon endothelial damage, platelets respond by spreading over the damaged area to prevent blood from leaving the circulation. We mimicked this injury in vitro by coating glass coverslips with fibrinogen, causing the platelets to attach to the surface, extend filapodia, and fully flatten out with lamellopodia formation. Representative pictures show that parthenolide substantially decreased the number of platelets able to fully spread onto a fibrinogen coated coverslip (Fig. 7A).

CD62P is a marker that is highly upregulated on activated platelets, assisting in transendothelial migration of leukocytes, thus inflammation [2]. While parthenolide treatment did not affect the basal percent of CD62P positive unstimulated platelets, it did decrease the percent of CD62P positive platelets following collagen activation (Fig. 7B). Soluble CD40L is a proinflammatory mediator abundantly released by activated platelets, and supernatant levels of platelet treatments were measured with ELISA. Parthenolide had no affect on basal secretion, but decreased soluble CD40L release when platelets were pretreated before collagen or thrombin activation (Fig. 7C).

In order to partially address the mechanism of parthenolide involvement in the altered activation of stimulated platelets, we assessed if oxidative stress alone could cause similar effects as parthenolide-pretreated platelets. Using H2O2 as a positive control,
we demonstrate that oxidative stress pretreatment of platelets before their stimulation with collagen did not affect the surface CD62P expression, and, in fact, increased the release of sCD40L (Fig. 7).

**Discussion**

Platelets are vital to hemostasis and have a critical role in immunological and inflammatory processes within human circulation. Severe thrombocytopenia often leads to hemorrhage, creating a rationale for developing thrombopoietic drugs. On the other hand, continuous activation of platelets is a major contributor to chronic inflammatory vascular diseases such as atherosclerosis and type-2 diabetes [2,28], creating the demand for new anti-platelet drug development. Either condition is detrimental, further exemplifying the delicate balance of adequate platelet numbers, and the risks of excessive platelet activation. We demonstrate here that parthenolide is a potential candidate agent for treatment of both conditions, as it increases platelet production from megakaryocytes and attenuates platelet activation during stimulation. Specific delivery mechanisms would need to be implemented, depending on the condition needed to be treated.

Two megakaryoblastic cell lines, Meg-01 and MO7e, can spontaneously produce platelet-like particles in culture [23]. We demonstrated that parthenolide facilitated morphological changes indicative of thrombopoiesis, and increased production of platelet-like particles within 24 hours of treatment (Fig. 1). Similarly, parthenolide enhanced platelet production within primary in vitro differentiated human megakaryocytes (Fig. 2). Compared to 15-deoxy-Δ12,14-Prostaglandin J2, which we previously reported as an enhancer of platelet production [4], parthenolide showed a weaker, but still significant enhancement of platelet production (comparison not shown). However, platelet production enhancement in a clinical setting by parthenolide and similar novel agents has not yet been assessed. It is worthy of noting that these primary megakaryocytes were first differentiated and matured with thrombopoietin (see Materials and Methods) before treatment of parthenolide. A bone marrow-directed conjunctive therapy may need to be considered before transition to an in vivo setting.

ROS and other oxidative stressors have been shown to increase after parthenolide treatment [5,7,11,29]. The increase of ROS in some cell types was associated with a decrease in GSH [5,29,30]. Contrary to those cell types,Meg-01 cells increased their total GSH levels (Fig. 3B), indicating they may be counteracting oxidative stress in this way, as the GSSG/GSH couple is considered to be the “cellular redox buffer” [26]. Meg-01 cells are capable of GSH depletion, as we demonstrated using BSO, but parthenolide does not act in this manner. Increase of GSH and HO-1, another antioxidant enzyme [24], along with no

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**Fig. 6.** Platelets derived from Parthenolide-treated primary human megakaryocytes are functional. PHM were treated with vehicle (veh) or 10 μM PTL for 24 hours and the produced platelets were then isolated and exposed to 10 ng/mL of collagen (Col) (1 h). Activation of the CD61 positive platelets was measured using an Accuri C6 flow cytometer. Compared to unactivated (top), collagen-activated platelet scatter plots (bottom) revealed an increase in granularity and decrease in size. PHM-derived platelets are activated similarly whether produced from vehicle or PTL-treated PHM (compare bottom two panels).
detectable ROS increase, suggests that the cells are capable of equilibrating to the induced oxidative stress caused by parthenolide. We showed that H$_2$O$_2$, which produces more ROS in megakaryocytes than parthenolide (Fig. 3C), and therefore generating an imbalanced redox state, was unable to affect platelet production from Meg-01, MO7e, or PHM cells (Fig. 4). Additionally, pretreatment of megakaryocytes with antioxidants before parthenolide treatment did not alter the increase of platelet production (data not shown). These data support the likelihood that oxidative stress does not play a major role in the enhancement of parthenolide-induced platelet production.

This led us to investigate another targeted pathway of parthenolide: NF-κB [10]. We have previously shown NF-κB signaling to play a role in megakaryocyte activation after PMA stimulation, and Zhang et al. showed that a constitutive NF-κB signal within proliferating megakaryoblasts decreased at final stages of maturation [13,19].

Using two different assays, we showed that parthenolide decreased NF-κB activity in megakaryocytes (Fig. 5). The rebound of NF-κB activity, as measured by luciferase at 24 hours, indicates that parthenolide is not toxic at this concentration to these cells, and also that the stability and duration of the effects of parthenolide is not altered. The more stable NF-κB inhibitor positive control, BMS, showed further reduction of luciferase activity. Additionally, this NF-κB inhibitor was also able to increase the number of platelets produced by megakaryocytes. While we cannot exclude the possibility that additional signaling pathways are also affected, we conclude that NF-κB inhibition is almost certainly a leading cause for the thrombopoietic effects of parthenolide observed in megakaryocytes. However, the short duration of parthenolide effectiveness may need to be addressed before transitioning into an in vivo model. More stable parthenolide analogs are already under investigation in the anti-cancer setting [31]. However, short lived efficacy of parthenolide may be advantageous if direct bone marrow delivery is feasible, as it may be harmful to give an anti-platelet drug intravenously to a thrombocytopenic patient.

We previously identified that the NF-κB pathway functions in normal human platelet activation [19], which is why we chose to investigate whether NF-κB inhibition within parthenolide-treated megakaryocytes affects the function of daughter platelets. Had this been the case, platelets created from parthenolide-treated megakaryocytes would have decreased activation ability. To the contrary, the platelets derived from parthenolide-treated megakaryocytes appear to be fully functional cells capable of activation (Fig. 6). This again, could be due to the short-lived effectiveness of parthenolide, therefore, only inhibiting NF-κB activity in mature megakaryocytes while it is not stable enough to affect daughter platelets. Importantly, this demonstrates that parthenolide enhances the production of functional platelets, making it a potential thrombopoietic drug candidate if it can be targeted to the bone marrow. Parthenolide is currently being investigated as an anticancer agent, but it may also have the potential to mitigate thrombocytopenia resulting from current therapies.

Anti-platelet drug therapies are increasingly sought as platelets are recognized as contributors to local and systemic inflammatory conditions such as atherosclerosis [2,28]. Anti-platelet therapies have also been shown to decrease platelets’ ability to mediate tumor cell invasiveness [32]. We found that when used as a pretreatment on platelets ex vivo, parthenolide decreased, but did not abolish the
activation of stimulated peripheral blood platelets [Fig. 7]. This evidence marks parthenolide as a potential drug candidate for anti-platelet therapy. Intravenous injections should be considered as a means of drug delivery for an in vivo setting, allowing parthenolide to quickly contact and dampen activated platelets before losing effectiveness.

Oxidative stress alone cannot be the mechanism of decreased platelet activation after parthenolide treatment because H2O2 did not affect platelet activation [Fig. 7]. We previously reported that NF-κB inhibitors can attenuate platelet activation [19], and our data suggest that the inhibition of this pathway by parthenolide led to the decrease in platelet activity reported here. Another proposed mechanism of the inhibition of this pathway by parthenolide led to the decrease in platelet activity reported here.

Our new evidence corroborates previously reported literature of parthenolide-driven reduction of platelet activity [16,17,35], and likely acts through multiple pathways including NF-κB inhibition.

Authorship contributions

J.S. designed and performed experiments, analyzed data, and wrote the manuscript. J.J.B., S.L.S., N.B., and R.P.P. all assisted in designing experiments and edited the manuscript.

Conflicts of interest statement

The authors declare no competing financial interests.

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