LAYERED, FLEXIBLE DRUG DELIVERY FILMS FOR THE PREVENTION OF FIBROTIC SCAR TISSUE FORMATION

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LAYERED, FLEXIBLE DRUG DELIVERY FILMS FOR THE PREVENTION OF FIBROTIC SCAR TISSUE FORMATION

DISSERTATION

A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy in the College of Engineering at the University of Kentucky

By
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ABSTRACT OF DISSERTATION

LAYERED, FLEXIBLE DRUG DELIVERY FILMS FOR THE PREVENTION OF FIBROTIC SCAR TISSUE FORMATION

Open wounds account for about 50% of military injuries and 10% of non-fatal traffic injuries. Scar tissue formation in these wounds may be reduced or prevented if treated with a combination of molecules whose release is tuned to the healing phases. The goal of this research was to develop flexible, layered drug delivery films for sequential, localized release of anti-inflammatory, anti-oxidant, and anti-fibrotic molecules to soft tissue.

Films were composed of cellulose acetate phthalate (CAP) and Pluronic F-127 (Pluronic). To impart flexibility, plasticizers, triethyl citrate (TEC) or tributyl citrate (TBC), were added. Mechanical analysis was performed on films as prepared and following phosphate-buffered saline incubation to determine property changes after implantation. Tensile tests revealed higher plasticizer content increased film elongation but decreased elastic modulus and ultimate tensile strength. TEC films elongated twice as much as those with TBC. After incubation, properties increased because plasticizer leached from films. Micro computerized tomography and scanning electron microscopy determined how erosion and plasticizer leaching affected the film’s structures before and after incubation. Porosity increased as plasticizer content increased; however, plasticizer content did not significantly affect erosion rates.

Next, effects of drugs with plasticizers on film erosion, release, and mechanical properties were investigated. Films were loaded with quercetin, an anti-oxidant, or pirfenidone, an anti-fibrotic, and plasticized with TEC or TBC. TEC-plasticized films containing quercetin released drug at a slower rate than TBC films. Pirfenidone-loaded films released drug at a faster rate than erosion occurred for both plasticizers. Increased pirfenidone loading resulted in significantly higher modulus and decreased elongation, an anti-plasticizer effect. Increasing quercetin loading significantly increased elongation. Size, solubility, and structure differences between quercetin and pirfenidone affected drug interaction with the films and the consequent mechanical and release properties.

Cell studies found TBC to be toxic even in low concentrations. Consequently, only TEC was further analyzed. Layered devices containing two drugs demonstrated
sequential release regardless of drug order. Plasticizer concentration did not significantly affect the release profiles. Lastly, in vitro and in vivo 9-layered device studies sequentially released drugs confirming the research objective: sequential, local release of anti-inflammatory, anti-oxidant, and anti-fibrotic molecules from CAP-Pluronic films.

KEYWORDS: Drug delivery, mechanical properties, plasticizer, cellulose acetate phthalate, Pluronic F-127.

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Open wounds accounted for 10.3% of the 5.3 million non-fatal injuries from traffic collisions in the U.S. in 2000, and about 50% of all military injuries involve musculoskeletal wounds to the extremities (1, 2). Chronic inflammation in these types of soft tissue defects can result in scars that have functional and aesthetic consequences. Fibrotic scar formation in these wounds may be reduced or prevented if treated with a combination of anti-inflammatory, anti-oxidant, and anti-fibrotic molecules targeted to the different healing phases. As such, the objective of this research was to develop and characterize flexible, layered drug delivery films for sequential, localized release of the aforementioned molecules to soft tissue.

Chapter 2, provides a review of the current state of the art on treatments for large soft tissue defects and a critical analysis of their shortcomings and potential areas of development. Scarless healing is examined and ongoing research in wound healing and drug delivery devices are discussed. To work towards developing a soft tissue wound healing device, mechanical and erosion properties of the system were evaluated (Chapter 3), drug release mechanisms were determined (Chapter 4), the cellular impact and capability of sequential release were assessed (Chapter 5), and finally in vivo studies were performed (Chapter 6). The films examined in this research are composed of two polymers, cellulose acetate phthalate (CAP) and Pluronic F-127 (Pluronic). Alone the
films are rigid. Since every wound has its own geometry and topography, the films need to be flexible and able to contour to the shape of any wound. To impart flexibility, plasticizers, triethyl citrate (TEC) or tributyl citrate (TBC), were added. In the third chapter, mechanical analysis was performed on plasticized films as prepared and following phosphate-buffered saline incubation to determine property changes after implantation. Micro computerized tomography and scanning electron microscopy determined how erosion and plasticizer leaching affected the film’s structures before and after incubation.

The combined effects of drugs with plasticizers on film erosion, release, and mechanical properties were investigated in Chapter 4. Films were loaded with quercetin, an anti-oxidant, or pirfenidone, an anti-fibrotic, and plasticized with TEC or TBC. Size, solubility, and structure differences between quercetin and pirfenidone were determined to affect how the drugs interacted with the films and the consequent mechanical and release properties.

In Chapter 5, cell studies were performed to evaluate the cytotoxicity of the TEC and TBC. The effect the plasticizers had on the mechanical properties of the films and the cytotoxicity results narrowed down the plasticizer choice for multi-layered devices. Two different multi-layered device types were evaluated, 4-layered devices and 9-layered devices. Both types demonstrated sequential drug release. Again, the differences in the drugs were found to affect the resulting release profiles. Pilot animal trials were performed to evaluate the devices in vivo and compare against in vitro
results in Chapter 6. Lastly, Chapter 7 is an assessment and conclusion of the developed and characterized devices against the objectives.
Chapter 2
Background and Significance

2.1 Clinical Need

With soft tissue defects accounting for over 10% wounds inflicted from motor vehicle accidents, over half a million car accident victims every year could benefit from wound healing films (1). Additionally, wounds caused during war, industrial accidents, or surgeries could also heal faster and with less scarring by the proposed drug delivery films.

2.2 Wound Healing

There are two categories of wound repair: primary unions and secondary unions (3). An example of a primary union wound is a surgical incision. The wound is narrow and clean and a limited number of epithelial and tissue cells die. Secondary unions are wounds in which a large mass of tissue has been destroyed and the wound cannot be neatly sewn together with sutures (4, 5). The healing steps are the same for secondary union and primary union wounds, but with secondary union wounds the process is longer, more complex, and the wound experiences more inflammation ultimately resulting in excessive scar tissue formation (6). Specifically, secondary union wound healing is the target of this research.
Wound healing is a complex process involving four main phases: hemostasis, inflammation, proliferation, and remodeling (7, 8). After the initial injury to tissues, the body undergoes hemostasis to stop blood loss (7, 9). During this phase, blood vessel walls have been damaged and platelets adhere to the collagen fibers present at the ruptured vessel walls. The platelets degranulate releasing adenosine diphosphate (ADP), calcium, and serotonin (10). These biomolecules or tissue damage initiates the plasma coagulation cascade. Incoming platelets exposed to ADP begin aggregating and start forming a platelet plug (11). The platelets degranulate further causing additional aggregation of platelets. Fibrinogen present in blood plasma is converted to fibrin which strengthens the clot.

Once blood loss is successfully controlled, the wound site undergoes inflammation (12). The signs of inflammation are redness, swelling, heat, pain, and sometimes loss of function. Preexisting chemical mediators, including histamine and serotonin, are released from local platelets and mast cells (3, 13, 14). These chemical mediators cause the surrounding vasculature to become leaky, predominately by endothelial cell retraction which causes a widening of the intercellular junctions (15, 16). The leaky vasculature allows an influx of fluid and cells into the injured tissue. Increased blood flow and fluid cause the previously mentioned signs of inflammation. Chemokines, chemotactic cytokines, are released by proximate cells and which causes recruitment of more cells to the area (17). The majority of the cells to enter the injured tissue are neutrophils and monocytes (10). They synthesize and release additional
chemical mediators, including arachidonic acid metabolites: prostaglandins and leukotrienes, cytokines, and nitric oxide (3). Nitric oxide and prostaglandins cause vasodilation allowing for the influx of more cells and extracellular fluid (18). The cytokines and leukotrienes activate other leukocytes to perform their inflammatory roles. Neutrophils protect against invading pathogens. Monocytes can differentiate into macrophages, which break down dead or invading cells and impurities through phagocytosis. Macrophages can congregate into one larger foreign body giant cell (FBGC) and remove larger debris (10). In normal healing, inflammation is a self-limiting process and the release of inflammatory mediators will reduce and cells will retract as the threat is resolved (19, 20). In chronic healing, the chemical mediators exacerbate the inflammation.

The next phase, the proliferation phase, replaces lost or damaged tissue. Angiogenesis occurs from surrounding blood vessels (21, 22). Macrophages present in the wound secrete epidermal growth factor (EGF) and vascular endothelial growth factor (VEGF). These cause endothelial cells in the blood vessels to bud and branch towards the wound, allowing for the exchange of nutrients and waste. Within a couple days, fibroblasts migrate to the injured tissue (23). The fibroblasts proliferate and create the proteoglycans and proteoglycan aggregates in the extracellular matrix (24). They repair the damaged tissue by forming a collagen matrix. If inflammation is not reduced, chemotaxis recruits excess fibroblasts to the newly formed granulation tissue. Growth factors, including platelet-derived growth factor (PDGF), fibroblast growth
factor (FGF), and transforming growth factor (TGF), cause fibroblasts to differentiate into proto-myofibroblasts (25-27). Then the proto-myofibroblasts differentiate into myofibroblasts which are responsible for expression of smooth muscle actin and create the contractile forces in the tissue (28).

Remodeling is the maturation of the scar tissue (3, 7, 12). This is the longest phase of all four since it can last for months. If inflammation continues during remodeling, cytokines and growth factors stimulate secretion of collagenases that breakdown the newly formed scar tissue (29). The collagen fibers will continue to be realigned, which can cause the contractile forces to become greater over time (30-32). These forces can cause disfiguration and malfunction around the injury site leading to the development of contracture, hypertrophic, or keloid scars (33-35).

2.3 Current Treatments

Secondary union wounds often contain bacteria (36). Either the object that severed the skin was unclean or the wound becomes susceptible to bacteria because of the large amount of exposed soft tissue (37). These wounds may also contain foreign materials. The first step to treating secondary union wounds is debridement, the process of removing foreign debris and dead tissue (6). There are five main types of debridement: surgical, mechanical, enzymatic, biosurgery, and autolytic (38, 39). In surgical debridement, necrotic tissue is surgically cut away and debris is removed with sterile tools. Saline irrigation, vacuum-assisted closure, ultra sound, and wound
redressing are types of mechanical debridement (39, 40). Pharmaceutically manufactured proteolytic enzymes are used applied to wounds in enzyme debridement. Collegenase is one of the most common enzymatic debridement methods because it selectively break downs triple helical collagen and disregards other proteins (29). In biosurgery debridement, steriley grown maggots are applied to the wound to remove necrotic tissue (39, 40). Autolytic debridement naturally occurs through cellular responses. Macrophages and FBGCs phagocytose debris and necrotic tissue (41). Phagocytosis occurs on the micron scale so some material can be too large or resistant for macrophages and FBGCs to break down (41, 42). If the debris is not removed a fibrous capsule will form around it (10). If a wound is infected, bacteria can create necrotic tissue faster than the macrophages and FBGCs can eliminate it. Depending on the severity of the wound, intravenous or oral antibiotics may also given prophylactically to kill any present bacteria (43).

Once the wound is cleaned, bandages are applied. Bandages are typically redressed every couple days (37). If a wound is too large for epithelium to fill in the defect, a skin graft may be required for the wound to close (6). There are two different categories of skin grafts: split-thickness skin grafts (STSG) and full-thickness skin grafts (FTSG) (44). Several factors must be considered before selecting a donor site, including contour, color, and vascularity. If a wound has not responded to traditional wound treatment, negative pressure wound therapy (NPWT) or vacuum-assisted closure
can also be used (45). NPWT creates a physical barrier, removes excess fluid, allows the wound to stay moist, and increases perfusion and granulation tissue formation (39).

Even though the treatment is more or less the same for all secondary union wounds, the healing time is not. Healing rates of chronic wounds are negatively affected by increased age of the individual wounded, increased size and decreased perfusion of the wound, and by reduced nutrition and health of the patient (3, 6, 46). Ischemia, diabetes, edema, and infection can also cause adverse healing (8). If inflammation continued through the remodeling phase, keloid, contracture, or hypertrophic scars are likely to develop (35). Once formed, treatments of these scars include surgical removal, cryotherapy, laser resurfacing, and steroid injections (47-50). If these treatments are utilized, the healing process of that location will occur again. Secondary union wounds often cause fibrotic scar tissue with aesthetic and functional consequences. There is much room and need for improvement upon current treatments.

2.4 Current Research

2.4.1 Scarless Wound Healing

During early development, human fetuses heal without any evidence of a scar (51-54). Fetal skin experiences a very minimal inflammation process during wound healing, which is related to the levels of cytokines and growth factors present (52). Transforming growth factors $\beta_1$ and $\beta_2$ have been shown to play a large role in the
development of a fibrotic scar experimentally. In fetal wounds, these are exhibited at very low levels (52). Interleukins (IL) 6 and 8 are found in low levels in fetal wounds and are much higher in adult wounds, while IL-10, a known anti-inflammatory biomolecule, is found in higher concentrations in fetuses than in adults. As discussed earlier, myofibroblasts mediate fibrotic tissue formation in adults; myofibroblasts are not even present in fetal flesh wounds until late gestation (52, 55). Knowledge gained through experiments with fetal wound healing can give insight into effective strategies to combat scar tissue formation in adult wounds.

2.4.2. Growth Factors and Wound Healing

As described above, the levels of growth factors present in a wound play a direct role in wound closure and scar formation. Hence, research is being conducted to better understand the effects of growth factors, including TGF, PDGF, FGF, and EGF, on wound healing (27, 56, 57). The effect of (EGF) on cutaneous scar formation in mice revealed that daily treatment, 100 µg, resulted in a third faster healing rate as compared to the saline treated control (58). The average scar size of the control group was 1.44 times as large as the average scar of the EGF treated group and the number of inflammatory cells present in the histological samples and the expression of TGF-β1 were much less for the EGF treated group (58). The concentration of EGF was found to be inversely proportional to the degree of scarring.
To study the effect of fibroblast growth factor (FGF) on wound healing, full-thickness dermal wounds were created in a rodent model (59). Chitosan films loaded with FGF were placed on the cutaneous wounds. There were three groups of experimental mice. One group’s films were loaded with 0.6 μg of FGF, another with 2.0 μg of FGF, and another with 6.0 μg of FGF. Another group of mice received an unloaded chitosan films on their wounds and the control group received only saline on their wounds. The FGF loaded chitosan films reduced the wound size and sped up the healing rate in a dose-dependent way (59). Histology revealed that the control group didn’t receive as high of an influx of macrophages and fibroblasts that the other groups did, nor did they experience as high a degree of angiogenesis in the healing wounds. In addition, FGF treated mice had more granulation tissue at the wound site as compared to the control.

Ischemic wounds often have adverse healing with excess inflammation (60, 61). PDGF and VEGF are growth factors that promote angiogenesis and correct an ischemic wound. Ischemia was inflicted on the hind limbs of mice (62). Immediately following surgery and again 5 days later, the experimental mice were treated with either PDGF, VEGF, or both PDGF and VEGF. Twenty one days after ischemia, the limbs treated with PDGF or both PDGF and VEGF had over 1.5 times the blood flow than the VEGF treated group or the control implying that PDGF is more important to ischemic wounds that VEGF (62).
2.4.3 Wound Healing Devices

Many different materials are being explored for drug delivery devices for wound healing applications including alginate (63-65), carboxymethylcellulose (66, 67), chitosan (68-71), collagen (72), dextran (73), hydrogels (65, 74-78), hydroxypropylmethylcellulose (HPMC) (79, 80), poly(vinyl alcohol) (PVA) (70, 81, 82), polycaprolactone (PCL) (83-85), poly(lactic-co-glycolic acid) (86-88), polyurethane (80, 89), and silk (86, 90, 91). Although all of these materials have been well studied for a variety of applications, very few are being evaluated as implantable wound healing devices. Instead, their effectiveness as topical dressings has been explored.

Porous gelatin and hyaluronic acid scaffolds were investigated as wound healing devices (92). When placed over split thickness wounds in a rat model, wound closure was accelerated and collagen deposition increased compared to the control which was no treatment. Electrospun wound dressings composed of PLGA and silk fibroin were shown to have a significant effect on wound closure in rats compared to PLGA dressings alone and the positive and negative controls after 15 days (86). Although the gelatin and hyaluronic scaffolds and PLGA fibroin wound dressings demonstrated an improvement to the control, they were replaced on the wounds every 24 and 48 hours, respectively, which is not possible with an implantable system.

Another challenge for implantable systems is coordinating the rates of drug release and erosion of the device. In topical wound dressings, that does not have to be considered. Films composed of electrospun PVA and PCL fibers and loaded with
caffeine, used as a model drug, were analyzed as potential flexible wound healing devices (93). These PVA and PCL films release drug on the order of hours and degrade on the order of weeks limiting the films to only dermal applications (84, 94). If these devices were implanted, they would leave behind material to cause an inflammatory response for far longer than the duration of the treatment (93). Hydrogels crosslinked with glutaraldehyde and loaded with gentamicin sulfate or norfloxacin were explored as an antimicrobial wound dressing (74). A full thickness dorsal wound was created in rats and the hydrogel was applied. Histology performed after 21 days revealed that the glutaraldehyde cross-linked hydrogels were found to have broken down into fragments and elicited an inflammatory response that interfered with the wound healing process (74) highlighting another challenge of the development of wound healing devices.

2.5 Proposed System

2.5.1 CAP-Pluronic

For over 20 years cellulose acetate phthalate combined with Pluronic F127 has been known to be a surface eroding drug delivery system (95). CAP and Pluronic can be easily fabricated into films, cylinders, or microspheres that can release a variety of different drugs (95-100). When loaded with drug in layers, devices were shown to release the drugs in a pulsatile manner (97, 100). Erosion and release rates of the system can be adjusted by changing the ratio of CAP:Pluronic; increasing Pluronic content increases the erosion and release rates while increasing CAP content decreases
the rates (95). The release mechanism from CAP-Pluronic devices is pH dependent. In neutral and basic conditions release is controlled through surface erosion and in acidic conditions, release is diffusion controlled (95). Previous studies have proven the system to be capable of releasing multiple drugs over several days, which is why it was chosen for this evaluation as a wound healing device.

2.5.2 Plasticizers

The CAP-Pluronic system alone is rigid. To allow a CAP-Pluronic film to contour to the varying geometries of a wound, plasticizers need to be added (101). There are several different families of plasticizers that are commonly used in medical and pharmaceutical applications (102-104). Glycols, citrates, sebacates in particular are frequently used to plasticize cellulose acetate (104). Phthalates were also commonly used to plasticize CAP, but in recent years, long term phthalate exposure has proven to have an adverse impact on health and development (105, 106). Triethyl citrate and tributyl citrate were chosen for the wound healing device developed because of their lower effect on erosion rates as compared to other plasticizers in addition to their biocompatibility.

2.5.3 Anti-inflammatory Drugs

There are two classes of anti-inflammatory drugs: steroidal and non-steroidal. Steroidal anti-inflammatory drugs typically require prescriptions and side effects can be
severe (107). Non-steroidal anti-inflammatory drugs (NSAIDs) act by inhibiting formation of prostaglandins by inhibiting the cyclooxygenase (COX) pathways that create them (108). There are two types of cyclooxygenase enzymes, COX-1 and COX-2. COX-1 is a common enzyme in healthy tissue and the inhibition of it can lead to gastrointestinal damage (109). Unselective NSAIDs inhibit prostaglandin synthesis by blocking both COX enzymes where selective NSAIDS preferentially inhibit either COX-1 or COX-2 enzymes (110). There are many different NSAIDs on the market including ibuprofen, acetaminophen, and aspirin. The drug delivery devices were evaluated in rodents first. NSAIDs approved for use in rodents include ketoprofen and meloxicam, so they were chosen for the device (108).

2.5.4 Anti-oxidant Drugs

There are two types of anti-oxidants: enzymatic and non-enzymatic. Of the non-enzymatic anti-oxidants, many different families exist: carotenoids, minerals, phenols, thiols, and vitamins (111, 112). They all reduce oxidative stress through a variety of mechanisms (113). Quercetin, a highly abundant dietary phenol, was chosen as the anti-oxidant drug for the device. Quercetin decreases superoxide activity of leukocytes and acts as a pro-inflammatory mediator (114-116).
2.5.5 Anti-fibrotic Drugs

Pirfenidone was the anti-fibrotic drug chosen for the wound healing film. Through several double blind studies, pirfenidone has proven to be an effective treatment for fibrotic diseases (117, 118). Pirfenidone reduces scar tissue formation by down-regulating cytokines and growth factors, including interleukin 1β and transforming growth factor-β1 (119-123). Other drugs, including imatinib mesylate and rosiglitazone, were developed as therapies for different conditions but inhibit pathways that create fibrosis. Imatinib mesylate inhibits growth factor receptors on mast and stem cells and rosiglitazone reduces inflammation that leads to fibrosis by reducing the inflammatory signal molecule, nuclear factor kappa-B (124-130).

2.6 Significance

Little research is currently being conducted on wound healing systems with the distinct aim of reducing fibrotic scar tissue formation. For an implantable drug delivery device to be effective at preventing scar tissue, it needs to be capable of altering the mechanisms of fibrotic tissue formation by providing therapeutic molecules over an extended period of time. The device also needs to erode as it delivers drug or be biologically inert, so the remaining vehicle does not perpetuate an inflammatory response (131, 132). Although research on topical wound dressings or treatments are promising, they require application of the therapeutic molecules on a daily basis over several weeks and cannot be implanted, only treat on one healing front by releasing
only one molecule, or leave behind a polymer vehicle in the wound that could cause adverse healing. There is a need for devices that are applied or implanted once, are capable of sequential release of multiple molecules to heal on multiple fronts, and erode as drug is released. This research was focused on the development and characterization of CAP-Pluronic films loaded with anti-inflammatory, anti-oxidant, and anti-fibrotic drugs for local delivery to soft tissue defects, with the ultimate goal of preventing or reducing fibrotic scar tissue formation and encouraging proper wound healing.

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Chapter 3

The Effect of Plasticizers on the Erosion and Mechanical Properties of Polymer Films


3.1 Introduction

While convenient, medications taken orally must make a long journey, traveling from the intestines, through the portal vein, into the liver, through the bloodstream until finally reaching their intended therapeutic target. During this convoluted process, many drug molecules are eliminated from circulation by the liver and kidneys, metabolized and excreted from the body (133-136). Because of this, oftentimes drugs must be given at doses far exceeding what the target site needs for treatment. This excessive concentration coupled with presence of the drug throughout all of the body systems results in a scenario where systemic side effects are guaranteed (137-139).

As a result, there is a need for a degradable, localized, drug delivery system capable of delivering drug to soft tissue so unnecessary sites are not medicated, reducing the risk of side effects, and drug is not wasted during clearance. Most of the current drug delivery systems are composed of slowly eroding polymer networks,
including Eudragit and poly(sebacic anhydride), from which drug diffuses out to provide therapy (140-145). This kind of device leaves behind material that may cause a foreign body reaction or cause inflammation from the byproducts as it slowly degrades. Also, many of the other currently researched systems are composed of rigid polymers, such as chitosan and Eudragit, that cannot contour to the shape of the treated site (145-147). The rigidity or stiffness of materials is related to their elastic modulus (148). Rigid polymers implanted in soft tissue will cause discomfort and possible fibrous capsule formation (10, 149, 150).

Plasticizers can be added to polymers to make them more flexible (102, 104). Plasticizers are small lubricating molecules that can change polymer properties and processability (104). In this study, the effect of plasticizers on the mechanical properties of an association polymer was explored. The system formed from cellulose acetate phthalate (CAP) and Pluronic F-127 has been proven to be a surface-eroding system that would release drug as the device eroded so no material would be left behind to cause an inflammatory reaction (95, 96, 151). Without plasticizer, however, the polymer is rigid and cannot be readily contoured. Two plasticizers, triethyl citrate (TEC) and tributyl citrate (TBC), have been included, and the system as a whole has been evaluated through mechanical analysis and degradation studies. Because these materials are ultimately intended to be implanted, their properties and structure when wet were also determined (152, 153).
3.2. Materials and Methods

3.2.1 Film Production

Films were made with a 70:30 weight percent ratio of cellulose acetate phthalate (CAP; Sigma-Aldrich, St. Louis, MO) and Pluronic F-127 (Sigma-Aldrich), respectively (Figure 3.1). Plasticizer, either tributyl citrate (Sigma-Aldrich) or triethyl citrate (Sigma-Aldrich), was added to the CAP-Pluronic mixture at either 0, 10, or 20 wt%. Acetone was added to make a 25% (w/v) solution. After vortexing the solution to ensure homogeneity, it was cast into Teflon dishes and then let stand in a 10°C refrigerator overnight. Films were desiccated overnight before testing.

Figure 3.1: Chemical structures of Pluronic F127 (a), cellulose acetate phthalate (b), triethyl citrate (c), and tributyl citrate (d).
3.2.2 Erosion Studies

Films were cut into discs using a circular punch measuring 5 mm in diameter. After determining their initial mass, samples were placed individually in a 24-well plate, and 1.85 ml of phosphate-buffered saline (PBS), pH 7.4, was added to each well. Plates were incubated on an orbital shaker plate 37°C. Half of the supernatant was replaced with fresh PBS every 12 hours. Three samples were collected and dried at different time points ranging from every 3 to 5 hours. Dried samples were weighed to determine final mass. The average mass loss was determined. The erosion rates were found by calculating the slope of average mass loss with time. Supernatants were collected from both TEC and TBC films after being incubated in PBS for 2 hours. The supernatants were pipetted into a 96-well plate and UV absorbance was measured at 630 nm using the BioTek PowerWave HT and Gen5 software to determine the opacity. PBS was used as a control.

3.2.3 Mechanical Properties

Films were cut to make microtensile test samples using a dog bone die (ASTM D 1708). Wet samples were incubated in 4 ml of PBS for two hours. Cross-sectional areas were measured for each film, both wet and dry, using calipers. Samples were mechanically tested to failure in tensile mode using the BOSE ELF 3300 with a ramp at a displacement rate of 0.5 mm/sec. Stress and strain were calculated, and elastic modulus (E), percent elongation, and ultimate tensile strength (UTS) were determined.
The percent elongation was normalized by the cross-sectional area so this value would not be influenced by thickness variations in the cast films.

### 3.2.4 Scanning Electron Microscopy

After dog bones were incubated in PBS for two hours, they were lyophilized and frozen at -80ºF for 30 min. Samples were then fractured so the cross-sectional area could be analyzed using a model S-3200-N Hitachi scanning electron microscope. The samples were coated with gold, and the microscope was operated at 20 kV and digital images were collected.

### 3.2.5 Microcomputerized Tomography (MicroCT)

To determine the interior porosity, microCT scans were performed using the SCANCO Medical AG µCT 40. Samples were cut to fit 12.3 mm tubes and were batch scanned. Scans were run in high resolution having 1000 projections with 2048 samples, a current of 177 µA, a potential of 70 kVp, and a 0º angle. Samples were contoured and evaluated with a program that distinguished between the polymer material and vacancies. The uniformly biased lower threshold was set to 20 so any part of the scan with a density below that was considered a void. The slices were reconstructed into a 3-dimensional representation of the scanned samples, from which the total volume, porosity volume, average pore size, and average spacing between pores were calculated.
using a morphometry script. Samples were scanned before and after being incubated in PBS for two hours.

### 3.2.6 Mass Spectroscopy

Mass spectra were acquired by the University of Kentucky Mass Spectrometry Facility. Supernatants from erosion studies were mixed an equal volume of acetonitrile before analysis. Electrospray ionization mass spectra were obtained on a ThermoFinnigan LTQ (ion trap mass spectrometer), with sample introduction by direct infusion at 3uL/min. Full scan mass spectra were recorded in positive ion mode. Instrument parameters included spray voltage: 3.5kV, capillary temperature: 185°C, capillary voltage: 50V, and tube lens voltage: 80V

### 3.2.7 Statistical analysis

Statistical analysis was performed using Instat (GraphPad Software). All of the samples with the same weight percentage of plasticizer or the same type of plasticizer were compared against each other. Results were analyzed using two-way ANOVA, and a p-value < 0.05 was considered statistically significant.
3.3 Results

3.3.1 Films

Plasticized films increased in flexibility as plasticizer content increased, while the unplasticized films remained rigid and could not be plastically deformed without failure (Figure 3.2). Qualitatively, films plasticized with TEC were more flexible than those plasticized with TBC at an equivalent weight percentage.

![Figure 3.2: A) Unplasticized inflexible CAP-Pluronic film and B) plasticized CAP-Pluronic film able to conform to any shape.](image)

3.3.2 Erosion Studies

Figure 3.3 shows typical mass loss curves for the control (unplasticized) and plasticized films. The addition of plasticizer did not significantly affect the erosion of the films. As the weight percentage of plasticizer increased, the magnitude of rate of erosion slightly increased from 2.2 wt% to 2.4 wt% per hour but the differences
between the three rates were not significant (Figure 3). As the films eroded, collecting the residual polymer to measure mass loss became a challenge so samples were only collected for the first 24 hours. The remaining mass of all of the films, plasticized with either TEC or TBC, after 16 hours was also statistically the same (Figure 3.4). Sixteen hours was chosen because at that point the initial mass loss due to plasticizer leaching had already occurred and the supernatant had been replenished with fresh PBS resulting in close to sink conditions. As TBC films eroded, the PBS became milky in appearance, whereas the supernatant remained transparent as TEC films eroded (Figure 3.5). Quantitatively, the opacity of the supernatant from the TBC sample was over 10 times as high as the TEC sample. The supernatant from the TEC sample was the same as PBS alone. Supernatants were analyzed by mass spectroscopy and after two hours the mass-to-charge ratios corresponding to TEC and TBC had a higher relative abundance than any other peak.
Figure 3.3: Example of typical degradation curves. Data are mean ± standard deviation (n=3).
Figure 3.4: Remaining mass after 16 hours of incubation in PBS at 37°C. Data are mean ± standard deviation (n=3). No differences were statistically significant.
Figure 3.5: Absorbance at 630 nm of supernatants from TEC and TBC films after 2 hours of incubation compared to PBS. Data are mean ± standard deviation (n=3). Inset: Images showing appearance of supernatants from TEC (above) and TBC (below) films.

3.3.3 Mechanical Properties as Prepared

Increasing the plasticizer content increased the effects on the mechanical properties; more plasticizer decreased the E and UTS and increased the percent elongation. Compared to unplasticized films, the 20 wt% samples plasticized with TEC had a lower E and UTS (Figures 3.6a and 3.6b), by about half, but had a 5 times higher percent elongation (Figure 3.6c). Addition of TEC to the films always significantly (p ≤ 0.05) affected the mechanical properties except between the ultimate tensile strength...
of the 10 wt% TEC and unplasticized samples. Adding 10 wt% TBC to the films decreased the UTS by roughly half, but addition of more TBC did not further affect the UTS. The 0 wt% TBC films had an elongation that was statistically the same as the 10 wt% TBC films, which had an elongation that was comparable to that of the 20 wt% TBC films. However, 0 and 20 wt% TBC films had elongations significantly (p ≤ 0.05) different from each other. The elastic modulus did significantly (p ≤ 0.001) decrease as more TBC was added to the films. For the concentrations examined, as plasticizer increased, the UTS of the TEC films decreased. With the TBC films, increasing plasticizer concentration past 10 wt% did not significantly affect the UTS. For similar plasticizer content, TEC caused greater changes to elongation, UTS, and E. For the 20 wt% samples, the TEC and TBC films had statistically similar UTS and E values, but their elongations were statistically different (p < 0.001), with and the 20 wt% TEC films plastically deforming over twice as much as the TBC films.
**Figure 3.6:** A) UTS of TBC and TEC films. 0 and 20 TEC were statistically different, \( p < 0.001 \). 0 and 20 TBC were statistically different, \( p < 0.01 \). B) Elastic modulus of TBC and TEC films. 0, 10, and 20 TBC were all statistically different from each other, \( p < 0.01 \). Both 10 and 20 TEC were statistically different from 0 TEC, \( p < 0.001 \) but 10 and 20 TEC were statistically the same. C) Elongation of TBC and TEC films normalized by the cross-sectional areas (CSA). 0 samples are statistically different from 20 TBC samples, \( p < 0.05 \). 0 was statistically different from 10 TEC, \( p < 0.05 \). 20 TEC was statistically different from both 0 and 10 TEC, \( p < 0.001 \). Data are mean ± standard deviation (n=3).
3.3.4 Mechanical Properties After Incubation

Allowing the films to soak in PBS for 2 hours drastically changed their properties, with the magnitude of some properties being five times different from those of the dry samples. With both TEC and TBC films, the UTS, E, and elongation (except for the 20 wt% TEC film) all increased after films soaked in PBS for 2 hours. When comparing the dry samples to their wet counterparts, the differences were always significant ($p < 0.001$) for all UTS data for both TEC and TBC samples (Figure 3.7a). The elongation significantly decreased for 0 and 10 wt% TEC samples when they became wet, but the 20 wt% had an elongation that was the same when wet or dry (Figure 3.7c). For the TEC samples, the elastic modulus was significantly ($p < 0.05$) affected by the duration in PBS for only the 0 wt% films (Figure 3.7b). Elastic moduli for the 0 and 20 wt% TBC films were significantly ($p < 0.05$) higher after incubation in PBS. For both TEC and TBC films, the elongation increased ($p < 0.001$) for 0 wt% and 10 wt% films, but not 20 wt%.
Figure 3.7: A) UTS of dry films and films incubated in PBS for 2 hours. Dry values compared to degraded values were all significantly different, p < 0.001. B) Elastic modulus of dry films and films incubated in PBS for 2 hours. 0 films were statistically different than their degraded counterparts, p < 0.05. 20 TBC was statically different from its wet counterpart, p < 0.05. C) Elongation of dry films and films incubated in PBS for 2 hours. Dry 0 and 10 TBC and TEC films were statistically different from their wet counterparts, p < 0.001. Data are mean ± standard deviation (n=3).

3.3.5 Morphology

Figure 3.8 shows representative SEM images of dry film surfaces and surfaces and cross-sectional areas after incubation in PBS for two hours. Samples appeared
smooth and had no porosity present before incubation. All of the samples had begun eroding after incubating in PBS for two hours. Porosity was seen both on the surfaces and in the cross-sections. As plasticizer content increased, more erosion occurred both on the exterior and interior of the film, resulting in increased porosity.
Figure 3.8: Scanning electron micrographs of the as-prepared surface (top left) and the surface (top right) and cross-section (bottom) of films incubated in PBS for 2 hours. A) 0, B) 10, and C) 20 wt% films.
MicroCT analysis quantitatively confirmed what was seen in the SEM images. Figure 3.9 shows cross-sectional slices of representative films before and after two hours in PBS, and the porosity of each film type is plotted in Figure 3.10. For TEC films, as plasticizer content increased, so did the porosity. The 20 wt% films increased to 17% porosity during incubation. The incubated samples were always statistically (p ≤ 0.05) different from their dry counterparts. Incubated 0 and 10 wt% samples were statistically the same, but both were significantly (p < 0.001) different than the porosity measured in the 20 wt% films. With TBC films, plasticizer content did not affect the porosity after 2 hours in PBS. Porosity increased uniformly to just under 1%. Pore sizes for both TEC and TBC films were found to be as small as 6 µm. The majority of the accessible volume in the films after erosion was from pores with a diameter of less than 20 µm. There were almost no pores present that were larger than 40 µm in either TBC or TBC plasticized films.
Figure 3.9: Micro CT images showing A) 0, B) 10, and C) 20 wt% films incubated in PBS for 2 hours.
Figure 3.10: Porosity measurements for as-prepared films and films incubated in PBS for 2 hours. A) TEC-containing films. 20 TEC was statistically different than 0 TEC, p
< 0.05. B) TBC-containing films. No statistical difference between incubated samples, and no statistical difference between non-incubated samples.

3.4 Discussion

CAP-Pluronic films are rigid and unable to conform to the varying topology of soft tissue defects that may occur in motor vehicle accidents or war wounds. In order to aid in placement of the films, imparting flexibility and allowing the films to contour to any shape, plasticizers were added. With the aim of making the FDA approval process quicker, plasticizers that were already deemed safe were chosen for the CAP-Pluronic films. Both triethyl citrate and tributyl citrate are common plasticizers used in pharmaceutical and biomedical devices (102, 103, 154, 155). The United States Pharmacoeopia deems both TEC and TBC appropriate to be used in pharmaceutical dosage forms (103). They are used in gelatin capsules, enteric coatings on pills, and transdermal drug delivery patches (154-156). They are also used in poly(vinyl chloride) and poly(vinyl acetate) components of medical devices, including tubing and films (157). Toxicology studies show that take orally, TEC can be toxic to rats at excessively high doses, corresponding to over half a liter in a 70 kg man. TBC is non-toxic in rats even at extremely high doses (158). For a wound that is 10 by 10 centimeters, a layered, drug loaded film would contain approximately 5 mL of plasticizer (0.07 mL/kg), which is well within the toxicity limits for a 70 mg man.

Triethyl citrate and tributyl citrate have molecular weights of 276.28 g/mol and 360.45 g/mol, respectively. This means that, for the same weight percent, more TEC
molecules are present in the films than TBC molecules (TEC: 0.74 mol per film for 10 wt%, TBC 0.58 mol per film for 10 wt%). The same molar concentration was not compared between TBC and TEC since plasticizers are most commonly added as a weight percent of the whole polymer. However, the 20 wt% films had 1.57x more plasticizing molecules than the 10 wt% TEC and they had very similar effects on the % elongation, UTS, and E, which shows that even with fewer molecules present, TEC plasticizes to a greater degree than TBC. Plasticizers with a lower molecular weight cause more flexibility in the material they are incorporated into because they have a higher mobility and allow polymer chains to more easily slide across one another (156).

With more lubricating plasticizer added to the films, a greater number of molecules were present between the polymer chains, which thereby increased the molecular separation and decreased the packing order of CAP and Pluronic F-127 (103). Because of this, for the same weight percent of plasticizer added, the TEC had more of an impact on the mechanical properties of the CAP-Pluronic F-127 films.

The mechanical properties changed rapidly after films were incubated in PBS, because much of the plasticizer leached out in the first two hours of degradation (159). Mass spectroscopy confirmed the plasticizer leach out by the presence of plasticizer in the supernatants of plasticized samples (results not shown). The TEC may leach out more quickly than the TBC because it is more hydrophilic than the TBC, and having smaller molecules also allowed it to travel down its chemical gradient faster. Triethyl citrate is a water soluble plasticizer up to 65 g/L, while TBC is a water-insoluble
plasticizer (160, 161). Differences in the solubility of TBC and TEC explain the differences in the supernatant’s appearance during degradation. The milky appearance was likely due to the TBC separating out from the PBS solution as the molecules leached out since it is insoluble in water, while the supernatant from the TEC films remained clear since it had not reached its solubility limit. Plasticizer leach out will not be an issue since the polymer will be in the shape of the wound and the film will only become stronger. After the film has been laid in place, there will no longer be a need for flexibility and the increased strength of the film will cause it to be a protective barrier during healing.

Pluronic F-127 may be eroding at a faster rate than CAP causing the porosity seen in the 0 wt% films. CAP is commonly used in enteric coatings because of its low water solubility. It dissolves at higher pH and therefore more slowly than Pluronic F-127 (162). This, in addition to the plasticizer leaching out, created the nanometer- and micron-sized voids seen in the SEM and microCT images. The pores increased the surface area and allowed the films to degrade at a faster rate. The films containing more plasticizer may allow the Pluronic degradation to occur more quickly. As voids are created in the film, PBS can penetrate the films more deeply. Water causes disassociation of the hydrogen bonds linking the ether oxygens from the Pluronic F-127 to the carboxylic acid groups on CAP, leaving behind a material that contains increasing amounts of CAP relative to all of the other materials. Because CAP is a stronger polymer than Pluronic and the association polymer they form together, erosion actually
increased the strength of the material (163, 164). The ultimate tensile strength and elastic modulus increased since the polymer chains could not stretch and begin to slide past each other as easily as before incubation. But interestingly, the elongation also increased. This is likely due to the small holes that were created in the polymer which allowed it to behave and stretch more like a sponge. As tension was applied to the wet material, the holes collapsed and caused the remaining chains to elongate. CAP is a stiffer, slower eroding material than Pluronic F-127, so even the 0 wt% samples had an increase in the elastic modulus and ultimate tensile strength as CAP became the predominant material in the films.

Before plasticization, the CAP-Pluronic system is less rigid than other drug delivery polymers such as Eudragit, which has a modulus of 500 MPa or chitosan, which has a modulus of over 1000 MPa without plasticizer (146, 165). CAP-Pluronic F-127 is a competitive system for drug delivery applications because it is surface eroding so drug is released as the device erodes, and since it erodes, no material is left behind to cause inflammation. Another beneficial property is its lower elastic modulus allowing it to be more flexible and contour and set to the shape of varying wounds when it is plasticized.

3.5 Conclusion

The degradation behavior and mechanical properties of CAP-Pluronic films can be varied by the type and amount of plasticizer incorporated into the system. The material properties will change shortly after the CAP-Pluronic drug delivery films are exposed to fluid. These changes do not affect the integrity of the system, since the
material only gets stronger and will maintain the shape formed in the tissue. The CAP-Pluronic system is attractive for a variety of soft tissue drug delivery applications because it could be tailored to have different properties.
Chapter 4

The Combined Effects of Drugs and Plasticizers on the Properties of Drug Delivery Films


4.1 Introduction

Large soft tissue defects are common in motor vehicle accidents as well as in military injuries. Open wounds accounted for 10.3% of the 5.3 million non-fatal injuries from traffic collisions in the U.S. in 2000, and about 50% of all military injuries involve musculoskeletal wounds to the extremities (1, 2). Aberrant wound healing results in extended inflammatory phases leading to the formation of fibrotic tissue (7, 12, 166). The greater the defect size, the longer the healing process will be (167), and infection and ischemia further prolong inflammation (60, 168). Wound location and the patient’s health and age also affect duration of the inflammatory phase (169). Secondary union wounds, where exposed soft tissue requires re-epithelialization, can result in scar contractures with negative esthetic and functional consequences (28, 58, 170). As such, there is a need for treatments that can simultaneously enhance wound healing while reducing fibrotic scar tissue formation.
Scar tissue formation could potentially be reduced or prevented if wounds were treated with individual or a combination of molecules targeted to the specific pathways of the healing process. The two drugs used in this study were quercetin and pirfenidone. Quercetin is a naturally occurring molecule that increases cell proliferation, decreases superoxide activity, and reduces wound contraction (113, 116, 145, 171). Reactive oxygen species are predominately released by infiltrating macrophages during the wound healing process and act as pro-inflammatory mediators (3). Thus, quercetin can inhibit oxidative stress to decrease inflammation. Pirfenidone is a pharmaceutical that has been shown to reduce fibrosis and scarring by down-regulating adhesion molecules and certain cytokines and growth factors, including interleukin 1β and transforming growth factor-β1 (119-123).

The wound healing device examined in this study was an erodible drug delivery film made of cellulose acetate phthalate (CAP) and Pluronic F-127 (Pluronic) (95). As the polymer erodes, drugs can be released from the system in a zero-order manner (96, 99, 151). CAP-Pluronic films are rigid, glassy polymers, but the addition of plasticizer increases flexibility, potentially allowing the devices to contour to the shape of varying wounds (101). The effects of plasticizers, in combination with different drugs, loaded in CAP-Pluronic films have not been previously studied. Films containing quercetin or pirfenidone with either triethyl citrate (TEC) or tributyl citrate (TBC) were investigated to determine the combined effects of drug and plasticizer on erosion, release, and mechanical properties.
4.2 Materials and Methods

4.2.1 Film Fabrication

Cellulose acetate phthalate (Sigma-Aldrich, St. Louis, MO) and Pluronic F-127 (Sigma-Aldrich) were combined in a 70:30 weight ratio, respectively, for a total mass of 2 g per film. Either tributyl citrate (Sigma-Aldrich) or triethyl citrate (Sigma-Aldrich) was combined with the CAP and Pluronic at 0, 10, or 20 wt%. Quercetin (10 or 100 mg; Sigma-Aldrich) or pirfenidone (6.1 or 61 mg; Tokyo Chemical Industries, Portland, OR) was added to the mixture (Figure 1). The low and high loadings of the two drugs are molar equivalents at 0.033 and 0.33 mol. A 25% (w/v) polymer solution was made by adding acetone to the CAP, Pluronic F-127, plasticizer, and drug. Mixtures were vortexed to ensure uniform dissolution of the components, cast into Teflon dishes, and the acetone allowed to evaporate at 10°C overnight. Films were desiccated overnight before analysis. Each sample is subsequently referenced by the drug loading within individual samples (approximately 64 µg, 640 µg, 39 µg, or 390 µg).

![Chemical structures of quercetin (a) and pirfenidone (b).](image)

**Figure 4.1:** Chemical structures of quercetin (a) and pirfenidone (b).
4.2.2 Erosion and Drug Release Studies

Five millimeter diameter discs were punched from the films. Samples were weighed to determine their initial mass and placed individually in 24-well plates. After adding phosphate-buffered saline (PBS), pH 7.4, to each well, the plates were gently agitated on an orbital shaker at 37°C. Three discs were collected and dried every hour, and their supernatants were saved for analysis of drug release. PBS in the remaining wells was replaced with fresh solution every hour. Dried samples were weighed to determine the final mass for calculating percentage mass loss.

High performance liquid chromatography (HPLC) analysis was performed using a Shimadzu Prominence system equipped with a Luna C-18 column (4.6 x 250 mm, 5 µm). For detection of quercetin, the mobile phase consisted of water containing 0.1% trifluoroacetic acid and methanol (30:70), with absorbance measurement at 254 nm. For detection of pirfenidone, the mobile phase consisted of water containing 0.2% acetic acid and acetonitrile (50:50), with detection at 310 nm. Injection volumes were 50 µl for all samples with a flow rate of 1 ml/min.

4.2.3 Mechanical Properties

Microtensile test samples were punched from the films using a dog bone die (ASTM D1708) and the width and thickness of each sample measured using digital calipers. Tensile tests were performed in displacement control mode at a rate of 0.5 mm/sec using a BOSE ELF 3300 system. From the force and displacement data, along
with the dimensions of each sample, the elastic modulus (E), percent elongation normalized by the cross-sectional area, and ultimate tensile strength (UTS) were calculated.

4.2.4 Statistics

Samples of the same drug molar content, drug type, same plasticizer concentration, and plasticizer type, were compared against each other. Results were analyzed using two-way ANOVA, and a p-value < 0.05 was considered statistically significant. Regression comparisons to a 45° line representing surface erosion were performed on slopes and intercepts of the release vs. erosion plots. A linear regression p-value < 0.05 was considered statistically significant.

4.3 Results
4.3.1 Erosion and Release Studies

The mass loss and drug release profiles for 0 and 10 wt% TEC and 0, 10, and 20 wt% TBC films loaded with 64 µg quercetin were linear (Figures 4.2a and 4.3a), with drug release rates from 9.3 to 13.0%/hr and erosion rates from 7.1 to 9.3%/hr. When mass loss and release were plotted against each other, quercetin-loaded TEC and TBC films followed along or below the 45° line that represents drug release based on only surface erosion (Figures 4.2b and 4.3b). The mass loss and drug release profiles for 20 wt% TEC films loaded with 64 µg quercetin both appeared more curved than the others but followed the 45° surface erosion line when plotted against each other. When the
same molar amount, 0.033 mol per film (39 µg per sample), of pirfenidone was added to the films, the cumulative drug release profile did not have the same linear behavior (Figures 4.2c and 4.3c). Over 9% more pirfenidone was released in the first six hours compared to the quercetin-loaded films (p < 0.001). None of the TBC-plasticized, quercetin-loaded films reached 80% cumulative release at six hours, but all of the TBC-plasticized pirfenidone-loaded samples had released over 93% of their total loading (Figures 4.3a, 4.3c, 4.3e, and 4.3g). Similarly, with the TEC-plasticized films, the cumulative release at six hours was under 83% for all of the quercetin-loaded samples and was over 92% for all pirfenidone-loaded samples (Figures 4.2a, 4.2c, 4.2e, and 4.2g). Pirfenidone release slowed significantly after six hours (Figures 4.2c and 4.3c). The TEC-plasticized films loaded with 39 µg pirfenidone released 98% or more of the drug at six hours, while the 64 µg quercetin-loaded films had a maximum cumulative release of 91% at that time (p < 0.001). When release from the TEC-plasticized films loaded with 39 µg pirfenidone was plotted against erosion, the first three data points were not significantly different from the 45° line for 10 and 20% TEC. For the last five data points, however, drug release was faster than the rate of mass loss (Figures 4.2d). The erosion versus release slope in this later region was significantly different from the 45° line (p < 0.01) for both 10 and 20% TEC. The plot of release versus erosion for the TBC-plasticized, 39 µg pirfenidone-loaded films was initially above the 45° line, and even after the first half of the mass loss, release still increased faster than the films eroded (Figure 4.3d). When the quercetin loading was increased to 640 µg, TBC films released
drug over 1.5 times faster than mass was lost and did not fit the erosion-based model (Figures 4.3e and 4.3f). TEC films loaded with quercetin released the drug at a significantly slower rate than did the TBC films: 52-56% of the quercetin had been released by three hours for TEC-plasticized films, while 74-80% had been released by TBC-plasticized films (p < 0.001). Quercetin release from TEC-plasticized films better followed the 45º erosion-based drug release line (Figures 4.2e and 4.2f). TEC and TBC films loaded with 39 and 390 µg pirfenidone released drug at a faster rate than erosion occurred (Figures 4.2f, 4.2h, 4.3f, and 4.3h).
**Figure 4.2:** Release and erosion of TEC films loaded with 10 mg quercetin (a), 6.1 mg pirfenidone (c), 100 mg quercetin (e), and 61 mg pirfenidone (g). Release versus erosion of TEC films loaded with 10 mg quercetin (b), 6.1 mg pirfenidone (d), 100 mg quercetin (f), 61 mg pirfenidone (h). Data are mean ± standard deviation (n=3).
Figure 4.3: Release and erosion of TBC films loaded with 10 mg quercetin (a), 6.1 mg pirfenidone (c), 100 mg quercetin (e), and 61 mg pirfenidone (g). Release versus erosion of TBC films loaded with 10 mg quercetin (b), 6.1 mg pirfenidone (d), 100 mg quercetin (f), 61 mg pirfenidone (h). Data are mean ± standard deviation (n=3).

4.3.2 Mechanical Properties

Higher plasticizer contents of both TEC and TBC increased the elongation and decreased the elastic modulus of CAP-Pluronic films (Figure 4.4). Increased pirfenidone loading, from 0 to 390 µg, in both TEC and TBC films caused a significant increase in E (p < 0.001) (Figure 4.4a and Table 4.1). TEC films with 20 wt% plasticizer showed a significant decrease in E when quercetin loading increased from 0 to 640 µg (p < 0.01) (Figure 4.4b). The modulus of films plasticized with TBC and loaded with quercetin increased with drug content from 0 to 64 µg and then decreased from 64 to 640 µg (p < 0.001 for 0 wt%; p < 0.01 for 10 and 20 wt%) (Figure 4.4b). Increasing pirfenidone loading from 0 to 39 µg significantly decreased elongation for all film types (p < 0.001) (Figure 4.4c). Quercetin-loaded samples had significantly greater elongation with increasing drug content (p < 0.001) (Figure 4.4d), and they elongated more than did pirfenidone-loaded films. For the same molar drug loading in 20 wt% TEC films, quercetin samples elongated 17 times more than did those containing pirfenidone (p < 0.001). For 20 wt% TBC, quercetin films elongated 26 times more than pirfenidone-loaded samples (p < .001).
Figure 4.4: Elastic modulus of pirfenidone-loaded films (a) and quercetin-loaded films (b). Percent elongation normalized by the cross-sectional area of pirfenidone loaded films (c) and quercetin loaded films (d). Data are mean (n=3).
Table 4.1: Standard deviations of (a) elastic modulus and (b) elongation for pirfenidone or quercetin loaded films.

a

<table>
<thead>
<tr>
<th>Plasticizer</th>
<th>TEC with pirfenidone</th>
<th>TBC with pirfenidone</th>
<th>TEC with quercetin</th>
<th>TBC with quercetin</th>
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<td>390µg</td>
<td>0µg</td>
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</table>

b

<table>
<thead>
<tr>
<th>Plasticizer</th>
<th>TEC with pirfenidone</th>
<th>TBC with pirfenidone</th>
<th>TEC with quercetin</th>
<th>TBC with quercetin</th>
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<tr>
<td></td>
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<tr>
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<td>0.12</td>
<td>0.11</td>
<td>0.03</td>
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</tbody>
</table>

4.4 Discussion

Two mechanisms govern drug release from CAP-Pluronic films: erosion and diffusion (172, 173). Erosion control release occurs when the loaded drug is released at the same rate at which the polymer vehicle erodes, which in the case of the CAP-Pluronic films is a zero-order process. In diffusion controlled release, water penetrates...
the material to dissolve the drug or disassociate it from the polymer, thus enabling drug to diffuse from the matrix. Erosion and diffusion control represent two ends of the spectrum; release from most degradable or erodible polymer systems fall somewhere between the two. The mechanism of drug release from unplasticized 70:30 CAP-Pluronic films has been shown to be surface erosion (95, 98, 99). Other drug delivery systems, such as poly(ε-caprolactone), poly(DL-lactide-co-glycolide), ethylene vinyl acetate, and starch, are diffusion-controlled and would leave behind polymer once release is complete (81, 85, 174-176). In contrast, CAP-Pluronic films release drug as they erode, so material will not remain in the healing wound to cause an inflammatory reaction after the therapeutic dose is released (10, 97). Polyanhydrides can also be surface-eroding systems but require stabilizing co-monomers increase hydrophobicity to slow hydrolysis and thereby drug release (177, 178).

The effects of plasticizers on release of drugs from the CAP-Pluronic system had not been previously studied. In other polymers, plasticizers can increase or decrease the rate of release by either increasing the surface area after plasticizer leaches out or by creating a better barrier to the dissolution media (179, 180). The plasticizers, TEC and TBC, incorporated into the films are already commonly used in pharmaceuticals and in biomedical devices (103, 104, 181). Triethyl citrate is a hydrophilic plasticizer with an aqueous solubility of 12 mg/ml, but TBC exhibits limited solubility at 0.15 mg/ml (182-184). Tributyl citrate is a larger molecule than TEC, with molar masses of 276.28 g/mol and 360.45 g/mol, respectively.
The two drugs explored, pirfenidone and quercetin, have different solubility in water, 4.4 and 0.36 mg/ml at 25 °C, respectively (185, 186). The molar mass of quercetin is 302.24 g/mol, which is over 1.5 times that of pirfenidone at 185.22 g/mol. Quercetin also has 12 bonding sites (5 donor and 7 acceptor) that can interact and/or interfere with the association polymer system. For example, the phenol and enol groups of the drug can form hydrogen bonds with the ether oxygens in Pluronic F-127 (186, 187). Additionally, the ketones in quercetin may interact with the carboxylic acid of CAP. These associations could affect polymer erosion as well as the release of quercetin from the system. In contrast, pirfenidone has only two acceptor sites on the amide that could bond with the CAP-Pluronic system (185). Fewer acceptor sites, a smaller size, and increased hydrophilicity resulted in pirfenidone being released more easily from the micron-sized pores created as the system eroded (101).

Samples containing pirfenidone exhibited a dual mechanism of release. Surface erosion controlled the first half of the 39 µg pirfenidone release from TEC-plasticized films. The first half of release from 39 µg loaded pirfenidone TBC-plasticized films also predominantly occurred via surface erosion, but after the fourth hour, release occurred at a faster rate than polymer mass loss, signifying a transition to diffusion controlled release. The shift in the main mechanism of release did not occur for the other drug and plasticizer combinations. The 64 µg quercetin-loaded films all fit the 45° line closely, indicating that the mechanism of release was surface erosion. If some diffusion occurred, the effect was insignificant. Previous work found that film porosity, even in
unplasticized films, increased significantly after a two hour incubation in PBS (101). Development of pores increased the surface area by which drug was released from the surface-eroding system. As more drug was added to the system, the leading mechanism of release shifted from erosion to diffusion and increased the release rate. This shift has been seen in other drug release systems, including atenolol loaded ethylene-vinyl acetate films (81). With increased drug loading, the concentration gradient is steeper and less CAP-Pluronic is present to control the release. The voids left behind by the released drug allow molecules deeper in the device to also diffuse out. This was seen for higher loadings of both quercetin and pirfenidone.

As observed in previous studies, increasing the plasticizer content of both TEC and TBC led to decreased elastic modulus and increased elongation, resulting in a CAP-Pluronic film that will conform better to the complex shapes of wounds (101, 188, 189). Triethyl citrate plasticized the films to a greater degree than did TBC because it is a smaller molecule, and for the same mass, there were over 1.25 times as many TEC as TBC molecules present in the films. Although pirfenidone is a smaller molecule than either of the two plasticizers, it still may have physically interfered with the ability of TEC or TBC to plasticize the films by separating the polymer chains and interacting with the carboxylic acid of CAP. In the unplasticized films, this pirfenidone interaction reduced what little flexibility the CAP-Pluronic films alone had, which resulted in an increased modulus and decreased elongation as pirfenidone content increased. It is not uncommon to see an “antiplasticizer” effect when small amounts of drug or plasticizer
are added to polymer (174, 188, 190). The chemical structures of pirfenidone and quercetin fit Jackson and Caldwell’s “antiplasticizer criteria” because they contain polar atoms, have at least two nonbridged rings, and have one dimension less than about 5.5 Å (190). The side groups of quercetin likely interacted with both polymers in the CAP-Pluronic system to result in synergistic effects on elongation and an increased modulus (191-193). The changes in the mechanical properties caused by the plasticizers and drugs did not correlate with differences in the release profiles. Mechanical testing occurred when the films were dry, and release occurred when the samples were wet, however. Previous work found that devices incubated in saline for two hours leached plasticizer resulting in increased strength (101). This phenomenon has been seen in other systems before, including sodium alginate-magnesium aluminum silicate films plasticized with polyethylene glycol 400 or glycerin (194). The chemical and mechanical changes that occur once the films are wet reduced any observable difference in the release.

4.5 Conclusion

Release from drug delivery films, particularly CAP-Pluronic films, can be controlled by the amount loaded and the drug properties, including size, hydrophobicity, and interactive side groups. The mechanical properties are also controlled by loading and the drug properties. Plasticizer can be introduced to further modulate films to achieve the desired mechanical properties. The combined effects of
drug and plasticizer on the properties of drug delivery films can range from antagonistic to synergistic. Different drugs and plasticizers can be added to CAP-Pluronic films to tailor the erosion, release, and mechanical properties needed for varying applications.
Chapter 5

Sequential Release of Multiple Drugs from Flexible Drug Delivery Films

This chapter will be submitted for publication, “Rabek C. L., Dziubla T.D., and Puleo D.A., Sequential Release of Multiple Drugs from Flexible Drug Delivery Films.”

5.1 Introduction

Sequential drug release is necessary for targeting the healing phases of soft tissue defects. After hemostasis, wounds undergo inflammation that, if not controlled, can result in excessive scar tissue formation during the proliferation stage (28, 195). Current research in drug delivery devices has fallen short in providing surface eroding systems capable of delivery multiple drugs in a sequential order and lasting for more than a couple days (196-199). In most systems, drug diffuses out leaving behind the polymer vehicle that could further inflame wounds.

The polymer films explored in this study were composed of cellulose acetate phthalate (CAP) and Pluronic F-127 (Pluronic). Together they form an association polymer that releases drug as the system erodes. Alone, CAP and Pluronic make a rigid polymer. To add flexibility to the system and allow it to contour to the shape of varying wound geometries, plasticizers were added. The two plasticized explored are citrates, triethyl citrate and tributyl citrate (Figure 5.2).
The drugs examined during these experiments were chosen to combat the scar tissue formation outlined in Figure 5.1. The anti-inflammatory, anti-oxidant, and anti-fibrotic drugs chosen were ketoprofen, quercetin, and pirfenidone, respectively (Figure 5.2). Ketoprofen is an anti-inflammatory drug that reduces inflammation by non-selectively inhibiting cyclooxygenase (COX) (63, 108). Quercetin was the anti-oxidant drug chosen because it decreases superoxide activity and reduces wound contraction (114-116). Pirfenidone, an anti-fibrotic drug, was chosen because it reduced scar formation by decreasing cell adhesion molecules and reducing cytokines and growth factors, including interleukin 1β and transforming growth factor-β1 (119, 121, 123).
Figure 5.2: Chemical structures of drugs (a) quercetin, (b) ketoprofen, and (c) pirfenidone, and plasticizers, (d) triethyl citrate and (e) tributyl citrate.

5.2 Materials and Methods

5.2.1 Cytotoxicity Studies

Mouse myoblast cells (C2C12; ATCC CRL-1772) were seeded into 24-well plates at density of 15,000 cells/cm² and cultured in DMEM supplemented with 10% fetal bovine serum at 37°C in a humidified, 5% CO₂ incubator. After 24 hours, the medium was changed, and cells were exposed to different concentrations of TEC or TBC plasticizer diluted into the DMEM media for 24 hours. To determine the remaining viability of the cells after plasticizer exposure, an MTT (3-(4, 5-dimethylthiazolyl-2)-2,5-
diphenyl tetrazolium bromide) assay was performed. Three hundred µL of MTT, 5 mg/mL, were added to each well and incubated for 2 hours. After replacing the medium with 500 µL of the extraction buffer, 20% sodium dodecyl sulfate in 50% N,N-dimethyl formamide, the plates were incubated at 37°C on an orbital shaker for 24 hr before reading absorbance at 570 nm.

5.2.2. Film and Device Fabrication

Polymer films were made by combining Pluronic F-127 (Sigma-Aldrich) and cellulose acetate phthalate (CAP; Sigma-Aldrich, St. Louis, MO) in a 30:70 weight ratio, respectively. Triethyl citrate (Sigma-Aldrich) or tributyl citrate (Sigma-Aldrich) were added to the CAP and Pluronic at 0, 10, or 20 wt% to create a 2 g film. Ketoprofen (100 mg; Sigma-Aldrich), quercetin (10 mg; Sigma-Aldrich) or pirfenidone (6.13 mg; Tokyo Chemical Industries, Portland, OR) were combined with the polymers and plasticizer. Acetone was added to make a 25% (w/v) mixture. To ensure homogeneity, the mixtures were vortexed. Then they were poured into Teflon dishes and kept in a 10°C refrigerator overnight to let the acetone evaporate slowly.

Two different device types were made, 4-layered devices and 9-layered devices. For the 4 layered devices, films were laminated using acetone and then punched into cylinders (6 mm). Ketoprofen and pirfenidone loaded films were separated by two blank films. The cylinders were inserted into polystyrene wells to ensure unidirectional erosion and release. Two device orientations were made, “forward” where the
ketoprofen layer was exposed first and “reverse” where the pirfenidone layer was exposed first (Figure 5.3).

![Diagram showing device types](image)

**Figure 5.3:** “Forward” (left) and “reverse” (right) device types.

For the 9 layered devices, the films were cut with cylindrical punches of increasing in diameter (Figure 5.4). The smallest layer (3.8 mm diameter), loaded with pirfenidone, was covered on either side by a blank layer of a larger diameter (5.5 mm diameter). The layers were laminated by painting acetone between the films. Larger diameter quercetin loaded layers (8 mm diameter) encased the blank layers. Another two blank layers (13.0 mm diameter) covered the quercetin-loaded layers. Ketoprofen-loaded films made up the outermost layers (18.2 mm diameter). All of the devices were desiccated overnight to remove all remnant acetone. Devices were then desiccated for 24 hours before analysis.
5.2.3 Drug Release Studies

Four-layered devices were placed individually into glass vials with 4 ml of phosphate-buffered saline (PBS). Samples were incubated at 37°C on an orbital shaker. Every 4 hours, supernatants were collected and replaced with fresh PBS. The nine-layered devices were placed into 50 mL test tubes with 30 mL of PBS. Every 12 hours, supernatants were collected (n=3) and replaced with fresh PBS. Supernatants were analyzed using high performance liquid chromatography (HPLC) to determine drug release at each time point. HPLC analysis was performed with a Hitachi Primaide system equipped with a Kinetix C-18 column (4.6 x 150 mm, 5 µm; Phenomenex). For detection of ketoprofen, the mobile phase consisted of water containing 0.1% TFA and acetonitrile (40:60), and UV absorbance was measured at 258 nm. For detection of pirfenidone, the mobile phase was water containing 0.2% acetic acid and acetonitrile (50:50) and
measurement occurred at 310 nm. Flow rates were 1 ml/min with injection volumes of 50 µl for all samples.

5.2.4 Erosion Studies

Initial masses were determined for the 9-layered devices. Every 8 hours devices were removed from the supernatant, lightly patted dry, and weighed.

5.2.5 Mechanical Properties

A dog bone-shaped die (ASTM D1708) was used to punch the four-layered microtensile test samples. Samples were desiccated overnight before calipers were used to measure the width and thickness of the laminated samples. Tensile testing was performed on a BOSE ELF 3300 system in ramp mode with a displacement rate of 0.5 mm/sec. The elastic modulus (E), percent elongation normalized by the cross-sectional area, and ultimate tensile strength (UTS) were calculated using the sample dimensions and force and displacement data.

5.2.6 Statistics

All of the samples of the same plasticizer concentration or plasticizer type were compared against each other. Results were analyzed using two-way ANOVA, and a p-value < 0.05 was considered statistically significant.
5.3 Results

5.3.1 Cytotoxicity Studies

Triethyl citrate was significantly less toxic to C2C12 cells than was tributyl citrate at 0.5, 1, 5, and 10 µL/mL (p < 0.001). Even at low concentrations, 0.5 µL/mL, TBC was found to be toxic to the cells (Figure 5.5). In contrast, TEC had no significant impact on cell viability at this concentration. At 10 µL/ml of TEC, only about half of the exposed cells were still viable. As the concentration of TEC increased, cell viability decreased. Because of these results, all further studies were performed using only TEC-plasticized films.

![Figure 5.5: Cell viability after plasticizer exposure. Data are mean ± standard deviation (n=3).](image-url)
5.3.2 Mechanical Properties

Increasing plasticizer concentration increased the elongation and decreased the ultimate tensile strength and modulus (Figure 5.6). Layered films that were not plasticized (0 wt % TEC) had a significantly higher elastic modulus and ultimate tensile strength than films that were plasticized ($p < 0.01$ and $p < 0.0001$, respectively). For elongation, every increase in plasticizer concentration was significantly different ($p < 0.05$).
Figure 5.6: Mechanical properties of 4-layered devices: (a) % elongation, (b) ultimate tensile strength, and (c) elastic modulus. Data are mean ± standard error (n=3).
5.3.3 Four-Layered Device Release Studies

For the 4-layered “forward” devices, peaks of ketoprofen and pirfenidone release occurred at 4 and 16 hours, respectively, regardless of plasticizer concentration (Figure 5.7). The overlap of the ketoprofen and pirfenidone peaks was 12 hours for all three plasticizer concentrations. The area under the two release curves corresponds to overlap release of both ketoprofen and pirfenidone. This “area under the curves” was also not significantly affected by the plasticizer concentration (Figure 5.8).
Figure 5.7: Release from “forward” 4-layered devices plasticized with (a) 0 TEC, (b) 10 TEC, and (c) 20 TEC. Data are mean ± standard error (n=3).
**Figure 5.8:** Area under the curves during ketoprofen and pirfenidone overlap release from 4-layered “forward” devices. Data are mean ± standard error (n=3).

For the “reverse” devices, the peak of the first drug, pirfenidone, still occurred at 4 hours (Figure 5.9). The peak of the second drug, ketoprofen, occurred at 24 hours for 0 wt% TEC devices, and at 16 hours for the plasticized devices. The overlap of the release of both drugs was 12 hours long for the 0 and 10 wt% TEC devices and 16 hours for the 20 wt% TEC devices. The area under the curve of the dual release was significantly different between the 20 wt% TEC devices and the 0 wt% and 10 wt% TEC devices (p <0.05) (Figure 5.10).
Figure 5.9: Release from “reverse” 4-layered devices plasticized with (a) 0 TEC, (b) 10 TEC, and (c) 20 TEC. Data are mean ± standard error (n=3).
Figure 5.10: Area under the curves during ketoprofen and pirfenidone overlap release from 4-layered “reverse” devices. Data are mean ± standard error (n=3).

5.3.4 Nine-Layered Device Release Studies

For the 9-layered devices, overlapping drug release was observed from the first time point with ketoprofen and quercetin (Figure 5.11). The ketoprofen peak is delayed with increasing plasticizer; peaks occurred at 8, 24, and 32 hours for 0, 10, and 20 wt% TEC, respectively. For all three device types, the peak quercetin release occurred at 32 hours. With increasing plasticizer concentration, the pirfenidone peak is release occurred earlier. The 0 wt% TEC devices had a pirfenidone peak at 144 hours, the 10 wt% TEC pirfenidone peak occurred at 84 hours, and the 20 wt% TEC devices at 64 hours. Pirfenidone is also more quickly released with increasing plasticizer. Pirfenidone
is first detected at 96, 64, and 32 hours for 0, 10, and 20 wt% TEC devices, respectively. Peak pirfenidone release for 0 wt% TEC samples was significantly different from the peak pirfenidone release from 10 and 20 wt% samples (p < 0.05 and p < 0.001, respectively).
Figure 5.11: Instantaneous release from 9-layered devices plasticized with (a) 0 TEC, (b) 10 TEC, and (c) 20 TEC. Cumulative release from 9-layered devices plasticized with (d) 0 TEC, (e) 10 TEC, and (f) 20 TEC. Data are mean ± standard error (n=3).

With increasing plasticizer content, increased mass gain occurred (Figure 5.12). Eight hours after the study began, the 20 wt% TEC devices gained 1.5 times their initial mass, the 10 wt% TEC increased in mass by 38%, and the 0 wt% TEC devices gained the least, with an additional 30% mass. The mass gain of the 20 wt% samples was significantly different from the 0 wt% samples (p < 0.01). The slopes of the mass loss
were 18.8, 21.2, and 25.0 for 0, 10, and 20 wt% TEC devices, respectively. The mass loss of the 20 wt% samples were significantly different from the 0 and 10 wt% samples (p < 0.001 and p < 0.05, respectively).

**Figure 5.12:** Mass loss of 9-layered devices. Data are mean ± standard error (n=3).

### 5.4 Discussion

Mouse myoblast cells were analyzed in the cytotoxicity study since they differentiate into muscle cells; muscle cells are what would be exposed to this polymeric drug delivery system. Tributyl citrate can be toxic to different organ systems, including the cardiovascular, respiratory, and central nervous systems (200-202). However, when taken orally, it is non-toxic even in high doses, equating 2 liters in a 70 kg man (158). Molecules delivered orally must travel from the intestines, through the hepatic portal
vein, and filter through the liver before circulating through the rest of the bloodstream. Many TBC molecules are excreted through the process and will never circulate through the body. The mouse myoblast cells examined did not have a lymphatic system or blood vessels to remove the TBC so they were unable to survive the toxic effects.

Compared to mechanical properties of single films plasticized with TEC or TBC (101), the four-layered films elongated more and had a lower UTS and modulus. Two of the four layers were drug loaded. The drug likely acted as plasticizer further affecting the mechanical properties of the films (191). Because of the additional thickness of the four-layered films, it is also possible that the acetone used to laminate the layers was not able to completely diffuse out of the polymer within 24 hours. Any remnant acetone would further plasticize the films (203).

Both the “forward” and “reverse” four-layered devices sequentially released pirfenidone and ketoprofen. The first drug released always had a sharp peak that occurred at 4 hours, whereas the second peak was much wider than the first, regardless of drug order. It was observed during the study that the center of the devices eroded faster than the outer edges, resulting in donut-shaped devices in the last few hours of release. A couple factors likely affected the second peak and the mechanism of erosion; the geometry and material of the wells. The geometry of the wells allowed for more turbulence towards the center of the wells; the walls of the wells protected the outer edges of the films. In addition to the geometry of the wells, the material they were composed of may also play a role in the inconsistent erosion. Polystyrene is a very
hydrophobic polymer, it may form nanobubbles on the surface preventing complete wetting of the material closest to the walls (204). This allowed the last drug-loaded layer to erode before the two blank layers separating them had completely eroded. Lastly, CAP-Pluronic films develop pores as they erode, and these pores could allow for diffusion of the second drug to extend its release peak (101).

The second drug release profiles in the 0 wt% TEC “forward” and “reverse” devices were quite different. The “forward” device had a steep slope before and after the peak release at 16 hours, while, the “reverse” device had a much gentler slope before and after the peak release. Differences in the drugs account for the release profiles. Pirfenidone and ketoprofen have different solubilities in water, 4.4 mg/mL and 51 µg/mL, respectively (185, 205). Ketoprofen is also 1.37 times larger than pirfenidone. Pirfenidone as the second drug was released more easily than ketoprofen. Ketoprofen, being larger and less water soluble than pirfenidone, cannot as easily diffuse out of the film as easily as pirfenidone. This is also supported by the differences between the “forward” and “reverse” overlap release. The “forward” 0 and 10 wt% TEC devices had larger overlaps of the two drugs than the “reverse” devices. Plasticizer did not affect release of pirfenidone as the second drug, but it did significantly increase the release of ketoprofen from 20 wt% TEC “reverse” devices compared to the 0 and 10 wt% TEC devices.

In the 9-layered devices, the concentration of TEC affected the release profiles of the three drugs. Increasing plasticizer increased mobility of the 3 drugs. TEC is 1.5x and
1.1x larger than pirfenidone and ketoprofen, respectively, and comparable in size to quercetin. As TEC leaches out of the films, it leaves behind pores large enough for the drugs to diffuse out (101, 206, 207). This increased porosity also accounts for the mass gain seen by the 10 and 20 wt% TEC films (208).

5.5 Conclusion

CAP-Pluronic films are surface eroding systems capable of sequentially releasing multiple drugs over several days. The release profiles can be tailored by the concentration of the plasticizer and by the type of drug loaded into the films. This makes them to be an appealing system for flexible drug delivery films, particularly for the prevention of scar tissue formation in soft tissue defects.
Chapter 6

Comparison of In vitro and In vivo Erosion and Release from Drug Delivery Films

6.1 Introduction

Chronic wound healing occurs when an injury undergoes excessive inflammation and often results in fibrotic scar tissue that can have both aesthetic and functional consequences (12, 209). In secondary union wounds, mass amounts of epithelial cells have been destroyed and the wound cannot be neatly sewn together with sutures. The healing process is longer and the wound experiences more inflammation than primary union wounds. Chronic inflammation causes the differentiation of superfluous myofibroblasts resulting in the development of fibrotic tissue (210).

Most common treatment for secondary union wounds begins with debridement, the process of removing foreign debris and dead tissue (6). Macrophages and foreign body giant cells will work hard to phagocytose debris and necrotic tissue. Once the wound has been cleaned, bandages will be applied and the wound will be redressed every couple days (37). During treatment, patients will also be administered oral anti-inflammatory drugs to help reduce pain and swelling. Depending on the severity of the wound, patients may take either non-steroidal anti-inflammatory drugs (NSAIDs) or corticosteroids. Current treatments do not target the pharmacotherapy for the different healing stages that occur. Also, with oral medications, much of the dose is lost
through the first-pass effect and what makes it to the bloodstream circulates through the whole body of the patient so most of the remaining molecules are provided to areas that do not need treatment. Hence, there is a need for localized treatments for soft tissue defects tailored to the healing process that can prevent or reduce fibrotic scar tissue formation and encourage proper wound healing.

Films composed of cellulose acetate phthalate (CAP) and Pluronic F-127 (Pluronic) and plasticized with triethyl citrate (TEC) have been proven to be surface eroding drug delivery systems capable of sequential release in vitro (95, 97, 99, 211). In this study, devices made from CAP-Pluronic films were loaded with anti-inflammatory, anti-oxidant, and anti-fibrotic drugs to determine the in vivo release profiles. The four drugs used were meloxicam or ketoprofen, quercetin, and pirfenidone (Figure A.1). Ketoprofen and meloxicam are common anti-inflammatory drugs that reduce inflammation by inhibiting cyclooxygenase (COX); ketoprofen is nonselective while meloxicam preferentially binds to COX-2 (63, 108). Quercetin is an anti-oxidant biomolecule that has been shown to increase cell proliferation, decrease superoxide activity, and reduce wound contraction when incorporated into a collagen matrix (114-116). Pirfenidone is an anti-fibrotic drug that has been shown to reduce scar formation by decreasing cell adhesion molecules and reducing cytokines and growth factors, including interleukin 1β and transforming growth factor-β1 (119, 121, 123).
Figure 6.1: Chemical structures of (a) meloxicam, (b) ketoprofen, (c) quercetin, and (d) pirfenidone.

6.2. Materials and Methods

6.2.1 CAP-Pluronic Film Fabrication

Cellulose acetate phthalate (Sigma-Aldrich, St. Louis, MO) and Pluronic F-127 (Sigma-Aldrich) were added in a 70:30 weight ratio to a test tube. Twenty weight percent triethyl citrate (Sigma-Aldrich) was combined to the mixture for plasticized devices. For drug loaded films, pirfenidone, quercetin, ketoprofen, or meloxicam were added to the 2 g mix of polymers at 6.1 mg, 10 mg, 100 mg, and 45 mg, respectively. Acetone was added to create a 25% (w/v) solution. To ensure homogeneity, the test
tubes were vortexed, then poured into Teflon dishes, and stored at 10ºC overnight. Acetone evaporation yielded casted films. Before device fabrication, films were desiccated overnight.

6.2.2 EVA Film Fabrication

Poly(ethylene-co-vinyl acetate) (poly(EVA)) 18 wt% vinyl acetate (Sigma-Aldrich, St. Louis, MO) was dissolved in 100 ºC toluene to make a 10% w/v solution. Once homogeneous, the solution was poured into a Teflon dish. Toluene evaporation left behind a cast film.

6.2.3 Device Fabrication

Cylindrical punches were used to make CAP-P layers increasing in diameter. The smallest layer, pirfenidone, was covered by the next smallest and so on until reaching the outermost drug loaded layer, ketoprofen or meloxicam (Table A.1). Layers were laminated with acetone. For the devices with backing layers, a poly(EVA) film was used to ensure unidirectional release. For devices with a mesh encasing, layers were covered in a polypropylene mesh that was stitched closed with nylon suture (Figure A.2). The devices were desiccated overnight to remove all remnant acetone. Devices were weighed and measured to determine their initial mass and dimensions.
### Table 6.1: Device fabrication information for animal trials

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of Devices/Rats</th>
<th>Layers</th>
<th>Backing Layer</th>
<th>Mesh Encasing</th>
<th>Drug Loaded Devices</th>
<th>Blank Devices</th>
<th>Plasticizer Types</th>
<th>Time Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>12</td>
<td>9 CAP-Pluronic Layers</td>
<td>No</td>
<td>No</td>
<td>Ketoprofen, Quercetin, Pirfenidone (6)</td>
<td>Yes (6)</td>
<td>0% and 20% TEC</td>
<td>48 and 72 hours</td>
</tr>
<tr>
<td>2</td>
<td>8</td>
<td>5 CAP-Pluronic Layers</td>
<td>Yes</td>
<td>No</td>
<td>Ketoprofen, Quercetin, Pirfenidone (4)</td>
<td>Yes (4)</td>
<td>20% TEC</td>
<td>24 and 48 hours</td>
</tr>
<tr>
<td>3</td>
<td>4</td>
<td>5 CAP-Pluronic Layers</td>
<td>Yes</td>
<td>Yes</td>
<td>Ketoprofen, Quercetin, Pirfenidone (4)</td>
<td>No (4)</td>
<td>20% TEC</td>
<td>24 and 48 hours</td>
</tr>
</tbody>
</table>
Table 6.2: Fabrication of devices used for the third trial

<table>
<thead>
<tr>
<th>Layer</th>
<th>Diameter (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polypropylene Mesh</td>
<td>19</td>
</tr>
<tr>
<td>Poly(EVA)</td>
<td>16.9</td>
</tr>
<tr>
<td>Pirfenidone</td>
<td>3.8</td>
</tr>
<tr>
<td>Blank</td>
<td>5.5</td>
</tr>
<tr>
<td>Quercetin</td>
<td>11</td>
</tr>
<tr>
<td>Blank</td>
<td>13</td>
</tr>
<tr>
<td>Meloxicam</td>
<td>16.9</td>
</tr>
<tr>
<td>Polypropylene Mesh</td>
<td>19</td>
</tr>
</tbody>
</table>

Figure 6.2: Eight layered devices for animal study.
6.2.4 Animal Methodology and Procedure

The animal study was conducted at the University of Kentucky in accordance with a protocol approved by the Institutional Animal Care and Use Committee (IACUC). The study consisted of three separate experiments: (1) devices without a backing layer or mesh encasement, (2) devices with a backing layer and no mesh encasement, (3) devices with both a backing layer and mesh encasement (Table A.2). Twelve week old, male Sprague-Dawley rats weighing between 350 and 400 g were used. The surgical area was prepared by shaving and disinfection with a povidone iodine solution. A roughly 3 cm incision was made on the left thigh over the quadriceps. Skin was separated from the muscle to create a pocket for the device. The device was placed into the pocket with the CAP-Pluronic side down (Figure 3). The incision was closed using wound clips. Rats were euthanized after 24, 48, or 72 h, and devices were retrieved. Final mass and dimensions of the devices were determined.

(a) 
(b)

Figure 6.3: Images of animal procedure: (a) implant in surgical site and (b) wound closure.
6.3 Results

All devices from the first animal study had completely eroded by the retrieval time points, 48 and 72 hours. Also, the animal study resulted in unexpected death for several of the rats implanted with drug loaded devices. The rats had an adverse reaction to the ketoprofen even though it is an approved and commonly used anti-inflammatory drug for rats. Rats that underwent planned euthanasia still had negative reactions to the ketoprofen, including porphyrin staining and diarrhea, and remained lethargic 48 hours after implantation.

For the second study, several parameters were modified. First the anti-inflammatory drug choice was changed to meloxicam, another approved anti-inflammatory drug for rats. To extend the device length in vivo, polypropylene mesh backing layers were bonded to the CAP-Pluronic layers to promote unidirectional release and erosion. Also, only 20% TEC plasticized devices were used, to reduce the number of rats needed and evaluate the worst-case-scenario in vivo duration devices (20% TEC devices were the fastest eroding devices). All of the rats lived until euthanized. When devices were retrieved at 24 hours after implantation, two of the four devices had completely eroded leaving behind only a backing layer. By 48 hours, all four devices had completely eroded.

Further modifications were made to extend the device life in vivo for the third study. A polypropylene mesh encased the CAP-P device and backing layer. For this study, four 20% TEC plasticized, drug loaded devices were used. Table A.3 shows that
devices used for the 48 hour time points were slightly larger, averaging an additional 0.024 g, or 7.5% more mass, than the devices used for the 24 hour time points. No blank devices were evaluated in this study. After 24 hours, the two devices lost 91.7% of their CAP-Pluronic mass (Figure 4). After 48 hours, the two other devices dropped in CAP-Pluronic mass by an additional 2.0%, with a total of 93.7% mass loss.

Table 6.3: Initial and final weights of \textit{in vivo} device components for third study.

<table>
<thead>
<tr>
<th>Timepoint (hours)</th>
<th>Initial Weight (g)</th>
<th>Initial Weight of CAP-Pluronic Layers (g)</th>
<th>Weight of Mesh (g)</th>
<th>Weight of Backing Layer (g)</th>
<th>Final Weight of CAP-Pluronic Layers (g)</th>
<th>Percentage of CAP-Pluronic Remaining</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>0.3165</td>
<td>0.2340</td>
<td>0.0700</td>
<td>0.0125</td>
<td>0.0188</td>
<td>8.0342</td>
</tr>
<tr>
<td>24</td>
<td>0.3178</td>
<td>0.2446</td>
<td>0.0643</td>
<td>0.0089</td>
<td>0.0208</td>
<td>8.5037</td>
</tr>
<tr>
<td>48</td>
<td>0.3259</td>
<td>0.2616</td>
<td>0.0717</td>
<td>0.0126</td>
<td>0.0092</td>
<td>3.8079</td>
</tr>
<tr>
<td>48</td>
<td>0.3558</td>
<td>0.2604</td>
<td>0.0737</td>
<td>0.0217</td>
<td>0.0228</td>
<td>8.7558</td>
</tr>
</tbody>
</table>
6.4 Discussion

Although ketoprofen has been considered a safe anti-inflammatory drug for use on rats, this study is not the first to observe adverse reactions. Shientag et al. found ketoprofen at 5 mg/kg doses to cause gastrointestinal bleeding and ulcers in rats after 24 hours (212). Combining anesthesia with the ketoprofen dose heightened the effects. Lamon et al. found the toxicity of ketorofen on Sprague-Dawley rats to be effected by their genetic strains and certain genotypes exhibited more exaggerated reactions (213). Ketoprofen is a nonselective cyclooxygenase (COX) inhibitor that interferes with prostaglandin production and inflammation reducing pain (108). Ketoprofen is known to irritate the intestine and kidneys resulting in ulcers. Thrombosis, by inhibition of
thromboxane $B_2$, will allow bleeding to continue from any consequential ulcers (108). Meloxicam is also an NSAID that is a cyclooxygenase inhibitor, preferentially binding to COX-2 (108). There is no literature supporting toxic interactions for meloxicam like are found with ketoprofen; COX-2 inhibitors are less likely to cause the gastrointestinal and renal damage that COX-1 inhibitors do, but they are also less effective as reducing inflammation (214).

The insignificant difference in erosion after 24 and 48 hours in the third animal study was caused by geometry changes over time, plasticizer leaching, and also possibly separation of the poly(EVA) from the CAP-Pluronic layers that was also seen in the second study. The ratio of the thickness to the diameter of the devices used in the third study was about 1:6. The device was thickest in the middle where all 8 layers were and got increasingly thinner towards the edges where fewer layers made up the thickness. CAP-Pluronic films are surface eroding systems (95). The surface area was its largest at $t=0$ resulting in the fastest erosion. As the outer edges eroded, the ratio of thickness to diameter increased and less relative mass was exposed on the surface, which slowed the erosion rate. The two devices used for the 48 hour time points weighed roughly 7.5% more but were no wider in diameter than the two 24 hour devices. This caused those two devices to have a lower relative mass exposure from $t=0$ which in turn resulted in slower erosion.

Most of the plasticizer leach out occurs in the first few hours of CAP-Pluronic erosion (101). Each CAP-Pluronic layer contained 20 wt% TEC. Because erosion
occurred so quickly, this leaching was captured in the mass loss of the 24 hour time points but not in the 48 hour time points. Plasticizer leaching has been shown to create porosity further increasing the surface area and increasing erosion. It is also possible that the poly(EVA) backing layer separated around the edges from the CAP-Pluronic layers allowing erosion to occur from both sides of the CAP-Pluronic films.

Compared to *in vitro* studies of layered devices, in Chapter 5, erosion occurred much faster *in vivo*. About 94% of the CAP-Pluronic in the *in vivo* devices was gone after 48 hours, whereas layered CAP-Pluronic devices without a backing layer and in sink conditions only lost 31% after 48 hours. This indicates that there are other factors increasing the erosion *in vivo*.

*In vitro* results cannot always predict what *in vivo* results will be (215). Chemical, enzymatic, and mechanical interactions can cause significant differences. Influx of inflammatory cells, including neutrophils and macrophages, to the wound may have sped up erosion through phagocytosis of the implant (9, 10, 216). Phagocytosis can occur for molecules on the micron scale; all of the components of this system, including the largest two, CAP and Pluronic, are that size or smaller (41, 42, 217). Enzymes *in vivo* may also have affected the erosion of the implant. The degree of substitution of cellulose acetate phthalate is 2.2 which prevents its cellulose backbone from being broken down into glucose units (218, 219). However, the ester bonds of the phthalic acid groups could be enzymatically broken. Since those functional groups are responsible for the association of the CAP and Pluronic, breaking its ester bond will
speed up erosion. In addition, mechanical erosion had a significant impact on the difference between the in vitro and in vivo results (220). CAP-Pluronic devices have been implanted on the calvaria of rats for other studies and significant mass loss after two days was not observed (97). The location of the implant had a significant effect on the erosion. The mechanical forces experienced by an implant are very difficult to model realistically in vitro. The rats in this study remained active and were not immobilized in any way. The rats activated their quadriceps anytime they needed to eat, drink, or play with their cage mate. The device was taut between the skin and muscle of the rat so as the muscle engaged, the devices were mechanically worn away. These mechanical forces could not be mimicked in vitro.

6.5 Conclusion

In vivo variances that cannot be replicated in vitro caused the CAP-Pluronic devices to erode faster than previously seen. Regardless, CAP-Pluronic devices still sequentially release drug and can be loaded with a multitude of molecules to locally treat an array of conditions. Furthermore, modifications could be made to prolong the duration of devices if needed, including adding slower degrading polymers between the layers, and allow for improved control over the drug delivery films.
Chapter 7

Conclusion

Improved treatments are needed for soft tissue defects to encourage proper wound healing and reduce scar tissue formation. In this research, a flexible, layered drug delivery film capable of sequential, localized release of anti-inflammatory, anti-oxidant, and anti-fibrotic molecules was developed and characterized to determine if it was feasible of meeting these needs. In the initial studies, mechanical analysis on plasticized films demonstrated that the properties and erosion of CAP-Pluronic films are varied by the type and amount of plasticizer incorporated into the system. The mechanical properties change shortly after exposure to fluid as plasticizer leaches out leaving behind a stronger film shaped by the surrounding tissue.

The following studies examined how characteristics of drugs, including size, hydrophobicity, and interactive side groups, influenced the release from CAP-Pluronic films. Mechanical properties of the films were also controlled by loading and the drug properties. Plasticizer can be introduced to further modulate films to achieve the desired mechanical properties. When drug and plasticizer were combined, the effect on the film’s properties ranged from antagonistic to synergistic. These studies concluded that different drugs and plasticizers can tailor the erosion, release, and mechanical properties needed to target healing in the soft tissue defects.
Layered CAP-Pluronic devices proved to be surface eroding systems capable of sequentially releasing anti-inflammatory, anti-oxidant, and anti-fibrotic molecules over several days. The devices maintained sequential release regardless of drug order.

Lastly, in vivo variances caused the CAP-Pluronic devices to erode faster than previously seen in vitro. Future work would involve reducing the erosion or mechanical wear. Changing the ratios of CAP to Pluronic, incorporation of a slower eroding polymer into the films, or modification of the implant site or animal model could also be explored to extend the device duration in vivo.

Current treatments for soft tissue defects are inadequate because they are either invasive, require multiple doses, or are not controlled to the healing phases to obtain optimal healing. CAP-Pluronic are a promising solution for the prevention of scar tissue formation in soft tissue defects because they can provide sequential release of anti-inflammatory, anti-oxidant, and anti-fibrotic drugs locally to healing wounds, which was the objective of this research.
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