Treatment for uterine fibroids: Searching for effective drug therapies

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Uterine fibroids are common reproductive-age benign tumors that contribute to severe morbidity and infertility. Cumulative incidence is 4 times higher in African-Americans compared to Caucasians and constitutes a major health disparity challenge. Fibroids are a leading indication for hysterectomy and their management averages $21 billion annually in the US. No long term minimally invasive therapies exist. Thus, promising drug therapies, their chemistry, pharmacology, and clinical efficacy, focusing first on innovative drug delivery approaches, are reviewed.

Introduction

Uterine fibroids are the most common benign pelvic tumor in women with a 70–80% cumulative incidence during childbearing years. African-Americans develop fibroids at younger ages than Caucasians and they tend to persist to menopause [1]. Fibroids tend to regress in size before menopause in Caucasian women. The etiology remains elusive although progress has been made. Previously reported similarities between fibroids and keloids [2,3] corroborate with recent findings [4]. Fibroids cells secret high levels of collagen and resist apoptosis. Ranging in location and size, growth is influenced by female gonadal steroids by apocrine and paracrine mechanisms. Subserosal or intramural fibroids can negatively impact fertility [5], and all fibroids are the leading cause of hysterectomy. Two Food and Drug Administration approved alternatives to hysterectomy are uterine artery embolization (UAE) and magnetic resonance-guided focused ultrasound surgery (MRgFUS) [6]. UAE is an angiographic technique that treats the whole uterus by causing ischemic fibroid necrosis. This therapy, like hysterectomy, is considered a standard treatment for women with no desire for future fertility. Alternatively, MRgFUS provides noninvasive fibroid-specific therapy utilizing high-intensity ultrasonography through the abdominal wall to cause coagulative necrosis in specific fibroids. Guidance and thermal monitoring is provided by dynamic real-time magnetic resonance imaging. However, fibroid regrowth after treatment in some cases may necessitate follow up therapies for both UAE and MRgFUS. Current studies are evaluating the long-term outcomes of these procedures. Thus, fibroids are a major public health challenge [7]. US treatment costs (~$21 billion annually) contribute more to healthcare expenditures than breast, colon or ovarian cancer [8]. As researchers develop next generation therapies, innovative approaches to drug delivery should be considered.

Drug therapies for uterine fibroids

Treatment options for fibroids vary as well as the severity of the symptoms, size and location of the fibroid lesions, the patient’s desire to maintain fertility, but the ultimate goal of

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therapy is the relief of the symptoms. As we learn more about the impact of fibroids on fertility, it becomes important for patients and their physicians to have a toolbox of therapeutic options with viable drug therapies being one of those tools. To this end, promising drug therapies, their chemistry, mechanism of action, pharmacology, clinical efficacy and side effects, focusing first on innovative drug delivery approaches, will be highlighted.

**Potential of novel therapies enabled by smart nanocarriers**

The development of nanocarriers designed to deliver and protect drug therapeutics (e.g. anti-fibrotic, aromatase inhibitors, progestins, among others) is an emerging field. Advances in guided-ultrasound technology (e.g. human *in vitro* fertilization where oocytes as small as 3–5 mm are manipulated) [9] make it feasible to envision utilizing nanocarriers to create a drug depot inside the fibroid by local injection. Thus, skilled physicians could inject the therapy into the uterine fibroid under guided-ultrasound in an outpatient setting. This approach would impede diffusion and distribution of the drug away from the injected fibroid, prolong release, delay inactivation, and therefore reduce the need for repeat injections. Examples of the most promising thermoresponsive delivery systems are given below.

**Atrigel®**

Atrigel® comprises a water-insoluble biodegradable polymer (e.g., poly(lactic-co-glycolic acid, PLGA) dissolved in a biocompatible, water-miscible organic solvent (e.g., N-methyl-2-pyrrolidone, NMP). A drug is added, forming a solution or suspension. Both the PLGA molecular weight and lactide:glycolide molar ratio governs drug delivery. Leuprolide acetate was incorporated into Atrigel® and depending on the targeted duration of leuprolide delivery, the L:G ratio in the PLGA varied from 50:50 to 85:15 and the polymer concentration varied from 34 to 50%. Clinical studies demonstrated a 22.5 mg leuprolide depot maintained an effective suppression of serum testosterone (50 ng/dL) for more than 3 months [10].

**ReGel®**

ReGel® is a ~4000 Da triblock copolymer formed from PLGA and polyethylene glycol (PEG, 1000 Da or 1450 Da) in repetitions of PLGA-PEG-PLGA or PEG-PLGA-PEG. ReGel® is formulated as a 23 wt% copolymer solution in aqueous media. A drug is added to the solution and upon temperature elevation to 37 °C the whole system gels. Degradation of ReGel® to final products of lactic acid, glycolic acid and PEG occurs over 1–6 weeks depending on copolymer molar composition. Chemically distinct drugs like porcine growth hormone and glucagon-like peptide-1 (GLP-1) may be incorporated, one at a time, and released from ReGel® [10].

**LiquoGel™**

We have engineered LiquoGel to work by mechanistically independent drug delivery routes: entrapment and covalent linkage [11]. This later feature distinguishes LiquoGel from other thermoresponsive injectables, as in theory, two or more drugs can be delivered to the tumor site. LiquoGel is a tetrameric copolymer of thermogelling N-isopropylacrylamide; biodegrading macromer of poly(lactic acid) and 2-hydroxyethyl methacrylate; hydrophilic acrylic acid (to maintain solubility of decomposition products); and multifunctional hyperbranched polyglycerol to covalently attach drugs. LiquoGel is formulated as a 16.9 wt% copolymer solution in aqueous media. The solution gels at physiological conditions and degrades to release drug contents within 1–6 days.

**GnRHa agonists and antagonists**

Gonadotrophin-releasing hormone analogues (GnRHa) target reduced production or hinderance of estrogen action on the smooth muscle cells of the uterus. Pituitary down-regulation suppresses the reproductive endocrine axis and can be exploited therapeutically to reduce circulating sex steroid levels. On the basis of this mechanism, several GnRH agonist peptides are commercially available in long-acting injectable depot formulations to suppress estrogen and thus create a menopause-like state. The agonistic response is produced by >-amino acid alterations in position 6 and/or ethyl-amide substitutions for carboxyl-terminal Gly10-amide (Fig. 1). These analogues are more resistant to proteolysis and increase GnRHa receptor binding affinity in the circulation, thus increasing the half-life of the analogues. As an example, triptorelin is used before surgery via injection for fibroid removal and, more recently, for preventative measures against permanent ovarian failure associated with chemotherapy [12].

By contrast, antagonists compete with endogenous GnRH for pituitary binding sites. Antagonistic analogues are formed by alterations in position 2 and/or 3. The first generation analogs were hydrophilic, and contained replacements for His at position 2 and for Trp at position 3. Third generation antagonists including cetrorelix and ganirelix were developed to eliminate anaphylactic reactions. One of the major remaining limitations to the wide use of the GnRH antagonists in leiomyoma treatment is the short half-life of these agents and the non-availability of depot formulations (necessitating repetitive dosing) [13]. Depot preparations of the next generation peptide antagonists degarelix and ozarelix are currently in late stage clinical development.

GnRHa treatment results in marked diminution of fibroid volume [14]. One mechanism of fibroid shrinkage by pharmacological concentrations of GnRHa was provided recently when decreased expression of nuclear factor of activated T cells 5 (NFAT5), a hyperosmolality gene expressed higher in
fibroid cells than myometrial cells, was noted [15]. This basal level expression is increased under hyperosmolar conditions, thus supporting the view that water flows out of fibroid cells at pharmacologic doses. GnRH agonists are widely used to prevent bleeding in preparation for surgical procedures in women with symptomatic fibroids. Long-term use increases risks of side effects including osteoporosis, vaginal dryness, impotence, reduced breast size, emotional instability, depression, hair loss, and musculoskeletal stiffness [7]. These side effects lessen with combinations of GnRHs and ‘add back’ therapies. Additionally, when GnRH agonist treatment by monthly depots is suspended, menses return after 8–12 weeks leading to a rapid increase of the uterus and fibroid size.

Aromatase inhibitors as a therapeutic drug strategy
Aromatase inhibitors (AIs) offer reversibility, total blockage of aromatase (CYP19), and possibly reduced side effects. AI medicines are often categorized as steroidal or non-steroidal aromatase inhibitors (NSAIs) based on their structural similarity with steroids, or as 1st and 2nd generations based on their evolution in time. Third generation AIs such as letrozole, anastrozole, and exemestane are used clinically. The half-lives are 48 h, 72 h, and 27 h respectively. Active research is ongoing to develop NSAIs that are more potent, selective and even multipotent [16]. Aromatase inhibitors are generally well tolerated with a low incidence of serious short-term adverse effects. Long-term use with the consequent hypo-estrogenaemia could result in loss of bone mineralization and an increased fracture risk.

The vast majority of women presenting with fibroid disease who would benefit from medical therapy are premenopausal, and aromatase inhibitors are unlikely to be effective. It should be noted, however, that in the obese menopausal woman presenting with fibroids, AIs may be preferable to progestin therapy as the latter has the potential to exacerbate the potential lipid disorder in the obese, often hypertensive woman.

Anastrozole and letrozole
This drug is associated with a reduction in fibroid size, thinning of endometrium and cessation of bleeding. Anastrozole has a half-life of ~48 h and is effective with daily oral administration. Mild side-effects of hot flashes, vaginal dryness and musculoskeletal pain are reported. The LEAP (letrozole, exemestane, and anastrozole pharmacodynamics) trial was a phase I pharmacodynamic study comparing the effects of these three AIs on safety parameters such as serum markers of bone formation and resorption, in healthy postmenopausal women with normal bone mineral density. The results demonstrated that all three inhibitors administered for 24 weeks caused incremental increases in bone resorption markers such as C-telopeptide crosslinks.

Selective estrogen receptor modulators (SERMs)
SERMs, like estrogen, are agents that elicit tissue-specific responses by intensely interacting with two kinds of estrogen receptors (ERs), ERα and ERβ, inhomogeneously distributed throughout the body. SERMs have complex pharmacokinetics properties due to their vaguely understood physico-chemical properties and low solubility in blood (1–200 ng/ml). In addition to analytical detection limitations at such low concentrations, several of the compounds have sufficiently long half lives that impede protocol development.
Since SERM are not administered to humans intravenously, the exact bioavailability of any of these drugs has not been evaluated properly. Frequently, differences in SERM activity depend upon the target gene promoter, as well as the background of a desired cell or tissue.

SERMs are characterized by their diverse range of agonist/antagonist actions on ER-mediated processes. SERMs belong to several different chemical classes such as benzopyran, benzothiophenes, chromane, indoles, naphtalenes, and triphenylethenes compounds. Many are available for clinical usage including raloxifene and tamoxifen discussed below. Novel SERMs are currently being tested in clinical trials such as LY353381 (arzoxifene), EM-652 and CP 336,156 and their structures are very similar to known SERMs [17]. The true breakthrough for SERMs could occur when ERα and ERβ are modulated independently with receptor-specific agents.

**Raloxifene and tamoxifen**

Two of the best characterized SERMs are tamoxifen and raloxifene, which are both considered to act predominantly as estrogen antagonists, blocking the effects of estrogens. Both raloxifene and 4-hydroxytamoxifen (a tamoxifen metabolite) fit into the hydrophobic pocket of the ligand binding domain of ER but the antiestrogen side chain prevents reorientation of helix 12 that must seal the ligand into the receptor before co-activators bind and produce a transcription complex. Both drugs interact through phenolic hydroxyls with Glu<sup>353</sup> and Arg<sup>394</sup> to correctly position the ligands in the binding domain. However, in the case of raloxifene, the side chain (crucial for antiestrogenic activity) interacts with Asp<sup>351</sup>. This same interaction is very weak in the case of 4-hydroxytamoxifen.

Raloxifene is a more complete uterine antagonist than tamoxifen, significantly reducing fibroid size in postmenopausal women yet is less efficacious at reducing tumor volume in premenopausal women [18]. Clinical outcomes in premenopausal women treated with raloxifene suggest that this compound, like tamoxifen, can affect the ovaries via the HPO axis. Tamoxifen is associated with insidious side effects, such as thromboembolic events, vasomotor symptoms and an increased risk of developing endometrial cancer and cataracts.

**Progestosterone receptor agonists and modulators**

The apparent, albeit unclear, role of progesterone (P4) in the growth of fibroids has encouraged the development of synthetic progesterone ligands with either progesterone receptor (PR) agonist (progestins) or mixed agonist/antagonist activity [19]. It has been shown that P4 and PR complexes reduce apoptosis and promote proliferation of fibroid cells [20]. PRs primarily exist in two isoforms, hPR-A and hPR-B. Both isoforms are acquired from the same gene by action of two different promoters [21]. Another PR isoform, PR-M has nongenomic activity and thought to participate in cellular respiration and protection from apoptosis. Recently, an increase in PR-M expression and mitochondria numbers was reported in fibroid edge compared to myometrium. Thus, a non-genomic progesterin-induced increase in cellular respiration may be a major factor in the growth of fibroids [22].

Figure 2 shows the chemical structure of various ligands that bind to PR ranging from full agonists on the PR to SPRMs. The progestins shown are derived from the parent molecule P4, although testosterone derived progestins also exit. Very small structural changes in the parent molecule may induce considerable differences in the activity of the derivative. Nonsteroidal SPRMs are actively being developed such as the 3-aryl indoles recently reported [23]. The alkyl substituent at N1 plays a very important role in binding potency and increasing the number of alkyl chains in this substituent group from methyl to ethyl or isopropyl (8) improved the binding affinity 7- and 5-fold, respectively. The functional activities follow this same trend.

Mifepristone is a 19-nortestosterone-derived compound that binds with high affinity to PR (slightly higher than P4) yet does not make the same binding contacts as progestins [24]. The side chains at the C11 position of the steroidal skeleton (R1 and R2) are the main structural features that correlate with the antagonist activity of mifepristone (Fig. 2). Two clinical trials of mifepristone reported a reduction in fibroid volume and uter size [24,25]. Side effects of mifepristone therapy include vasomotor symptoms and hot flashes (increased over baseline in the 10-mg group). Although promising, patients experienced slow regrowth of fibroid tumors following cessation of drug treatment.

Levonorgestrel is a hormonally active levorotatory enantiomer of a racemic mixture of norgestrel, a progestin derived from 19-nortestosterone. The drug produces an antiproliferative action on the endometrium. Levonorgestrel is formulated within an intravaginal device (IUS) and commercially sold as Minerva<sup>®</sup>. The device is inserted directly into the uterus and its use is attributed to significant reduction of uterine bleeding [26,27]. While uterine volume decreased, there is controversy as to whether or not significant fibroid shrinkage occurs [28]. There is a high expulsion rate when the IUS device is used for the treatment of heavy menstrual bleeding associated with fibroids (9.8%) or in the presence of submucous fibroids (15.4%) [28].

Asoprisnil is a hydrophobic oxime with substitutions at the C11 position of its steroid-like skeleton. Clinical trials involving asoprisnil have reported reversible suppression of menstruation and variable effects on ovulation (Phase I) [29]; reduced fibroid size in a dose-dependent manner (5, 10 and 25 mg), and a similar effect on improvement with menorrhagia with a 28–83% amenorrhea rate (Phase II) [29]; improved dysfunctional bleeding and reduce fibroid
size (Phase III) [19]. These results are again promising, however, SPRMs like asoprisnil have not yet received FDA approval due to concerns over the effect of these agents on the endometrium.

Ulipristal (5) and its acetate analog (6) were developed as SPRMs with pure PR antagonistic activity. They both have minimal antiglucocorticoid effects and are under development for use in symptomatic fibroids. In vitro studies and small randomized trials involving ulipristal report fibroid size and related symptoms were significantly reduced ($P = 0.01$ and $P = 0.04$) over placebo with no adverse effects observed [30]. Ulipristal acetate treatment administered for 13 weeks on randomly controlled women with symptomatic fibroids, menometrorrhagia, and anemia, successfully controlled excessive bleeding due to uterine fibroids and reduced the size of the fibroids [31]. Changes in total fibroid volume calculated by MRI have also been used as primary outcome measure for success of ulipristal therapy [32]. Ulipristal acetate is currently in a Phase III clinical trial.

Telepristone has mostly progesterone antagonist activity and low antiglucocorticoid activity. It was specifically developed for use as an antiprogestin in the treatment of uterine fibroids and endometriosis. It has been shown to reduce fibroid size by up to 40% and significantly decreases vaginal
bleeding. While one study reported that telepristone induced apoptosis in fibroid cells [33] another studies did not confirm this fact [34,35]. The US FDA placed a full clinical hold on telepristone in August 2009 because of elevated liver enzymes associated with drug treatment. Since the partial clinical hold status granted in June 2010, one clinical trial with low-dose telepristone designed to test the safety of the drug and efficacy of escalating doses has been initiated.

**Chemical inhibition of growth factors**

TGFβ is the main profibrotic cytokine which regulates fibrosis by stimulating activation, survival and collagen synthesis of collagen producing cells. This dimeric polypeptide, composed of identical 112-amino acid subunits, is over expressed in uterine fibroids compared with myometrium from the same individual [2]. Possible therapies could develop from TGFβ neutralizing antibodies, soluble analogues of TGFβ receptor to act as a decoy receptor, or orally administered TGFβ receptor inhibitors [36]. However, targeting TGFβ is problematic, requiring proteins as therapeutics (cost, purity, stability, immune response) and manipulating TGFβ signaling inhibition, preventing its widespread usage.

**Anti-fibrotics**

Fibroids are characterized by altered collagen fibrils, fibrosis and tissue stiffness [37,38], as well as increased amounts of type I and V collagen [39]. Selective elimination of collagen producing cells or reducing their state of activation is currently limited to experimental trials. Medical strategies that interfere with collagen formation (e.g. anti-fibrotic drugs) should be efficacious in fibroid treatment. A high throughput screening assay has been developed to indentify drug candidates that show antifibrotic activity [40]. Two promising antifibrotic candidates are highlighted below, but in general this class of pharmaceuticals presents undesirable side effects when systemically administered.

**Pirfenidone**

Pirfenidone is an orally bioavailable antifibrotic agent that has been shown to regulate fibrosis [41]. Pirfenidone inhibits fibroblast proliferation, diminishes the messenger RNA levels of collagen types I and III in a dose dependent manner, and effectively inhibits myometrial and fibroid cell proliferation in vitro [42]. The effect of this antifibrotic drug in fibroid cells is specific only for collagen type I. The drug is currently in phase III clinical trials for the treatment of pulmonary fibrosis [43].

Pirfenidone is relatively insoluble in aqueous solutions (<2%) but very soluble in alcohol and chloroform. This drug penetrates the cell membrane without requiring a receptor. When administered orally, Pirfen is easily absorbed in the gastrointestinal (GI) tract, reaching most tissues and crossing the blood–brain barrier. Pirfenidone displays rapid absorption ($t_{\text{max}} = 0.33–1$ h) and clearance ($t_{1/2} = 2–2.5$ h) [41]. Two metabolites were identified, which seem to be produced from oxidation of the methyl group on the pyrrolidone ring followed by subsequent formation of the carboxylic acid. Oral administration of pirfenidone in clinical trials is associated with undesirable side effects including vomiting, fever, abnormality of hepatic function, dizziness, facial paralysis, hepatoma, and skin photosensitivity.

**Halofuginone**

As an extract from hydrangeas, halofuginone is a small organic molecule exhibiting coccidiatostat benefits in birds and more recently anti-fibrotic activity against fibroid cells. Halofuginone inhibits both fibroid and myometrial smooth muscle cell proliferation by rapidly inhibiting DNA synthesis and later inducing apoptosis [44]. In addition, halofuginone significantly suppressed TGFβI mRNA production.

After single intravenous and subcutaneous administration, halofuginone rapidly distributes out of the plasma compartment and into the tissues resulting in a large volume of distribution. Renal clearance was limited to 7.5–16.7% of total body clearance and only 15–16% of the parent compound was excreted in the urine within 48 h after oral administration. Plasma-half life varies between 5 and 17 h. At 3.5 mg/day, nausea, vomiting, and fatigue were reported. Several patients experienced bleeding complications on treatment with halofuginone in which a causal relationship could not be excluded. The PKs of halofuginone were linear over the dose range studied with a large interpatient variability. This medication is not currently used in humans and its toxicity is unknown [45]. Reported side effects of halofuginone when taken in patients with advanced solid tumors were nausea, vomiting and fatigue.

**Purified collagenase**

Collagenase clostridium histolyticum is a FDA approved drug targeting Dupuytren’s Contracture in adults with a palpable cord. This drug comprises a fixed-ratio mixture of two classes of purified collagenases; clostridial type I and type II collagenase. The type I enzyme is 114 kDa 1000 amino acid single polypeptide chain and type II collagenase is about 1000 amino acids long with a molecular weight of 113 kDa. Both are metalloproteinases requiring zinc and calcium for full activity and have selective activity against collagen. These two classes of enzymes are not immunologically cross-reactive and differ from each other in domain structure substrate affinity and catalytic efficiency. When combined, they demonstrate synergistic collagenolytic activity.

Class I collagenase enzymes attach to the C and N termini of the mature triple helix collagen while class II enzymes attach to bonds within the collagen molecule. For catalytic activity, zinc is required and calcium is necessary to maintain the conformation of the collagen binding sites in a state
that allows the enzyme to bind to the triple helical collagen fibril. Both classes are capable of degrading small collagen fragments. The concerted effort of both enzymes degrades the entire collagen triple helix as opposed to mammalian matrix metalloproteinases which degrade only specific bonds in the collagen molecule and do not degrade collagen completely. In vivo, clostridial histolytic collagenase is not effective in degrading type IV collagen. Thus in clinical studies, purified clostridial histolytic collagenase did not degrade large blood vessel membranes or nerves.

In mammals, including humans, clostridial histolytic collagenase is inhibited rapidly in the blood stream by serum proteins which forms a complex with the collagenase. These complexes containing the inactive enzyme are then removed from circulation in the liver. Purified clostridial collagenase is not a reproductive toxin.

The dose used for treatment of a Dupuytren’s contracted joint is 0.58 mg per direct injection into the cord. Up to three injections may be given per affected joint. This drug is approved by FDA for the treatment of Dupuytren’s Contracture of the adult hand with a palpable cord present and has not yet been used in clinical trials for fibroids. Our laboratory is conducting proof of principle studies.8

Low-dose oral contraceptive pill as a therapeutic drug strategy

Combined oral contraceptive pills and progestin-only pills are effective for the treatment of abnormal uterine bleeding [46]. These therapies can induce endometrial atrophy and stabilize the endometrium. However, the myoma size does not change. Further, evidence shows that estrogens and progestins can act as growth stimulators for uterine myomas.

Herbals as a therapeutic drug strategy

Herbal therapies are recognized in Eurasia for their benefits to human well being. Traditional Chinese Medicines (TCM) are documented for modern pharmacological characteristics in a recent book chapter [47]. Kue-chin-fu-ling-man (KBG) improved clinical symptoms of hypermenorrhea and dysmenorrhea in 110 premenopausal women with uterine fibroids and fibroid shrinkage was observed in ~60% of the women. TCM relate fibroid formation to stagnated blood (‘yu zheng’) accumulation. Thus, blood circulation enhancers are often added to the herbal formulation.

Other Chinese formulations (Augmented Rambling Powder, Cinnamon Twig and Poria Pill) are reported to lower estrogen levels [48]. Dong Quai and Peony Powder reduce blood estrogen levels and inflammatory compounds in the uterus. Four Substance Decoction treats both fibroids and endometriosis in women who are affected by poor diet or overfatigue [48]. Western herbal remedies are also highlighted [48]. Cinnamon oil (5–10 drops) administered 16 times in 4 h is reported to subside bleeding due to fibroids. Reishi stops pelvic inflammation when administered daily in triplicate dosages of 12 mL/dose. Green tea is a powerful antioxidant touted as a remedy for various symptoms including fibroids [13]. Approximately 90–126 mg dry weight of the active ingredient, catechins, is typically contained in 1 g leaf brewed for 3 min in 100 mL water. Epigallocatechin-3-gallate (EGCG) is the major component of catechin (>40%) of the total polyphenolic mixture of green tea catechins. The mechanism of EGCG action appears to be blockage of tumorgenesis by modulating signaling pathways involved in cell proliferation, transformation, inflammation, and oxidative stress. All these processes contribute to the pathogenesis of uterine fibroids. EGCG induced apoptosis and inhibited proliferation of fibroid cells [13]. This suggests that EGCG may be a potential anti-fibrotic agent acting through multiple signal transduction pathways. An ongoing double-blind placebo-controlled clinical trial is currently in phase II of investigations to evaluate the clinical role of EGCG in women with symptomatic fibroids [13].

Conclusions

If the molecular basis for fibroid development and of myometrial proliferation is understood, additional nonsurgical therapeutic interventions may be forthcoming. Current clinical needs are stated to a) include an effective prevention strategy if possible b) improve early detection; c) slow the growth of fibroids; d) determine the mechanisms of infertility due to fibroids; e) develop better treatment modalities; f) reduce recurrences after treatment; g) evaluate long-term results.

Drug therapies for uterine fibroids are either short-lived or have significant side effects. Leuprolide acetate is presently used for only short-term treatment before surgery. SPRMs have not yet received FDA approval, due to concern over the effect of these agents on the endometrium, despite the intensive developmental efforts by several pharmaceutical companies and clinical trials which demonstrated significant shrinkage of fibroids and symptomatic improvement. Other medical therapies have been suggested in the recent past such as SERMs, AI’s, and anti-fibrotics but clinical trial results for efficacy treating fibroids has been limited or disappointing. TGFβ inhibitors effectively inhibit fibrosis in different animal models; however, systemic inhibition of TGFβ raises important safety issues because of the pleiotropic physiological effects of this factor. Local delivery of such agents allows the delivery of intact peptide molecules and promises to reduce fibrosis, while avoiding systemic side effects.

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Questions arise as the knowledge about uterine fibroids continues to develop. Patients and their care givers will wonder: Is there a drug therapy that rivals the effectiveness of surgical procedures yet preserves the uterine child bearing function? If realized, could such a therapy be administered under ultrasound guidance in an outpatient setting? Addressing these questions presents unique opportunities at the interface of molecular medicine and clinical care.

With the advent of the means to deliver drugs or drug combinations directly to fibroid tumors, the potential of reduction and perhaps eradication of fibroids before the need for surgical intervention could be realized. Multiple drugs could be given with the LiquoGel platform as a drug cocktail or sequentially for the benefit of patient care. Several more conventional drug therapies for uterine fibroids could be potentially entrapped or covalently linked to LiquoGel to afford delivery with potentially reduced side effects, improved efficacy, and controlled release profiles. The treatment could be administered by skilled personnel in a doctor’s office under ultrasound guidance.

Ultimately, development in uterine fibroid therapy stems from the need for viable choices to reduce symptoms associated with this disease. In light of rising health care costs, patients looking for economical approaches to health may consider a combination of conventional medicine supplemental approaches. Both care giver and patient should be educated about herbal treatment options for uterine fibroids. Although further studies and chemical investigations are needed to determine and validate the herbal therapy efficacy for uterine fibroid treatment, patient interest in herbal treatments may excite more investigation in this field.

Conflict of interest
The authors have no conflict of interest to declare.

Acknowledgements
This work was supported by National Institute of Health K12HD043446. NIH was not involved with any aspects of preparing this manuscript. We would like to thank Friederike Jayes and Aletheia Burrell for assistance with developing this review.

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