Application of Bioactive Hydrogels for Functional Treatment of Intrauterine Adhesion

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Intrauterine adhesion (IUA) is a common endometrial disease and one of the main causes of infertility in women of childbearing age. Current treatment strategies, such as hysteroscopic adhesion resection, hysteroscopic transcervical resection of adhesion (TCRA), the use of local hormone drugs, and anti-adhesion scaffold implantation, do not provide a satisfactory pregnancy outcome for moderate-severe IUA, which presents a great challenge in reproductive medicine. With the development of material engineering, various bioactive and functional hydrogels have been developed using natural and synthetic biomaterials. These hydrogels are not only used as barely physical barriers but are also designed as vectors of hormone drugs, growth factors, and stem cells. These characteristics give bioactive hydrogels potentially important roles in the prevention and treatment of IUA. However, there is still no systematic review or consensus on the current advances and future research direction in this field. Herein, we review recent advances in bioactive hydrogels as physical anti-adhesion barriers, in situ drug delivery systems, and 3D cell delivery and culture systems for seeded cells in IUA treatment. In addition, current limitations and future perspectives are presented for further research guidance, which may provide a comprehensive understanding of the application of bioactive hydrogels in intrauterine adhesion treatment.

Keywords: bioactive hydrogel, reproductive medicine, drug delivery system, stem cell therapy, intrauterine adhesion

INTRODUCTION

The uterus is composed of three tissue layers, namely, the endometrium, myometrium, and perimetrium (Mancini and Pensabene, 2019). The endometrium is the innermost layer composed of epithelial and stromal components; it is a unique tissue that undergoes a repetitive cycle of cell proliferation, differentiation, and shedding during the reproductive years of a woman’s life, providing the “fertile ground” for embryo implantation (Lv et al., 2021). Intrauterine adhesion (IUA) is a result of endometrial injury and infection caused by unsafe abortion and poor maternal care, which can lead to partial or complete occlusion of the uterine cavity, resulting in periodic abdominal pain, oligomenorrhea, amenorrhea, even infertility (Liao et al., 2021). A normal endometrium is a determinant of fertility, and IUA is recorded to be the second most common cause of female infertility after fallopian tube obstruction (Wei et al., 2020). As for IUA treatment, current therapies include hysteroscopic adhesiolysis, TCRA, hormonal therapy, and anti-adhesion material implantation, including the intrauterine device (IUD) (Huang et al., 2020a; Han et al., 2020). Although current treatment strategies have achieved positive effects in some cases, the prognosis of
patients with severe IUA is unsatisfactory, and the incidence of re-adhesion after the operation is still high (Young and Evans-Hoeker, 2014). Thus, patients always accept adjuvant therapy after operation to avoid the recurrence of re-adhesion after surgery (Li et al., 2021). To date, clinicians have already applied biomaterials such as hyaluronic acid (HA) and INTERCEED—an absorbable adhesion barrier made of oxidized regenerated cellulose—after surgery in IUA patients to prevent postoperative re-adhesion (Zhang et al., 2020a). In addition, numerous adjuvant therapies and biomaterials, including bioactive hydrogels, can be selected for those patients who suffer from severe IUA. With the rapid development of regenerative medicine and tissue engineering, in addition to applying traditional adjuvant therapy to IUA patients, clinicians are also seeking the potential of endometrial regeneration through tissue engineering.

Hydrogels are formed by water-soluble or hydrophilic polymers through certain chemical or physical cross-linking, and are composed of a hydrophilic three-dimensional (3D) network structure. The hydrogels swell rapidly in water and can retain a large volume of water without dissolving in this swollen state, which is very similar to soft tissue (Milcovich et al., 2017; Manna et al., 2019). Hydrogels are widely applied in the field of tissue engineering owing to their excellent properties in drug release, 3D cell culture, and simulation of an extracellular matrix (Chang et al., 2018). In the treatment of IUA, the bioactive hydrogels show potential in anti-adhesion and endometrial reconstruction. In order to improve the biocompatibility, reduce the potential cytotoxicity, and adapt to the special pathological microenvironment of the IUA, researchers have further constructed and optimized hydrogels (Tang et al., 2018). Although hydrogels have performed outstandingly in tissue regeneration, cell culture, and drug delivery, their application in the field of reproductive medicine is still in the exploratory stage. Only a limited number of hydrogel products and technologies have been successfully used in humans (Huang and Ding, 2019).

In this review, we summarize the latest advances in hydrogels as a therapeutic option for IUA. This article focuses on the latest research regarding hydrogels as physical anti-adhesion barriers, in situ drug delivery systems, and 3D cell delivery and culture systems, in order to improve their application in IUA (Scheme 1).

**THE DESIGN OF HYDROGELS FOR IUA**

The design of hydrogels for biomedical applications needs to consider the purpose of the application and the disease-specific microenvironment. IUA can be characterized as endometrial fibrosis, where the endometrium is surrounded by a muscular layer of interlacing smooth muscle fibers, which usually causes the uterine cavity to collapse during the postoperative healing process unless it is mechanically distended. Since the uterine cavity is varied in size and shape, the bioactive hydrogels which can adjust themselves to different uterine cavities may fully promote endometrial regeneration. Moreover, the speed of degradation of the hydrogels needs to be manageable to avoid rapid degradation because they must remain in the uterine cavity for a certain time to resist adhesion recurrence (Wang et al., 2020a). After the healing process is completed, the bioactive hydrogels as barriers must be degraded or absorbed naturally, rather than remaining as a foreign body. Among the hydrogel systems sensitive to stimuli,
thermosensitive hydrogels are the most extensively studied. Depending on the properties, thermosensitive matrices are divided into systems showing lower critical solution temperature (LCST) or upper critical solution temperature (UCST) (Kasiński et al., 2020). In this field, temperature-responsive hydrogels have attracted the attention of researchers; especially the hydrogels that are liquid at room temperature and can be rapidly gelled at physiological temperature around specific tissues (Kapoor and Kundu, 2016). The most important hydrogels for biomedical applications are systems with LCST close to the physiological temperature. This type of hydrogel is widely used in anti-adhesion materials because of the ease of its control and its rapid response to physical and chemical changes (Keskin et al., 2019). Compared to other hydrogels, temperature-responsive hydrogels can mechanically help shape the normal uterine cavity and prevent postoperative IUA (Han et al., 2016).

Secondly, hydrogels applied in IUA treatment should have controllable release profiles. As a simple and efficient platform for controlled release and delivery, the ideal bioactive hydrogels embed growth factors, drugs, and stem cells with therapeutic functions within their 3D structure. These drugs are released through the pores, hydrolysis of bonds, or self-degradation of the hydrogels, allowing the drugs to be applied to the uterine cavity wound after surgery with a sustained profile to promote regeneration of the endometrium (Sheth et al., 2019). Chemically crosslinked hydrogels usually provide a stable polymer network with slow degradation kinetics. For patients with severe IUA, where there is almost no normal functional endometrial tissue in the uterine cavity, stem cell transplantation is a potential option for the promotion of endometrial regeneration. Under such circumstances, the ideal bioactive hydrogels need to create a 3D cell delivery and culture system that is conducive to transplanted stem cell survival, proliferation, and directed differentiation (Cervelló et al., 2015).

Depending on the source, hydrogels can be divided into natural hydrogels and synthetic hydrogels. Natural hydrogels (e.g., HA, alginate, chitosan, collagen, and polyethylene glycol) have already been widely used in the field of regenerative medicine (Liu et al., 2020a). They are generally highly biocompatible, as reflected in their successful use in the peritoneum and other sites in vivo (T et al., 2016). Their biocompatibility is promoted by a high-water content and a physiochemical similarity to the native extracellular matrix, both compositionally (particularly in the case of carbohydrate-based hydrogels) and mechanically (Larrañeta et al., 2018). The naturally derived materials often have desirable biological properties and can influence cell function, but some natural hydrogels are limited by poor mechanical strength and fast degradation profiles (Rastogi and Kandasubramanian, 2019). In contrast, synthetic polymers provide appropriate 3D environments and have the desired mechanical strengths. However, they lack the bioactive properties of natural materials. Therefore, it is necessary to produce hybrid materials by combining synthetic and natural polymers, and to retain the desirable characteristics of both materials (Liu et al., 2018).

To date, hydrogels used in clinical settings can only achieve physical anti-adhesion. Bioactive hydrogels with controlled release and suitable culture microenvironment have not been widely used in the treatment of IUA. Hydrogels have been widely exploited with the aim of achieving drug delivery directly into the vaginal mucosa for hormone therapy, owing to their attractive features such as the increased drug residence time at the targeted sites and control of drug release rates after hysteroscopic transcervical resection of adhesion (TCRA) (Shamloo et al., 2018).

THE APPLICATION OF BIOACTIVE HYDROGELS IN IUA

The excellent biocompatibility, controllable mechanical properties, the ability of incorporating therapeutic agents for sustained release, and encapsulating seed cells at physiological conditions make bioactive hydrogel an ideal option for IUA treatment and endometrial regeneration. Several factors should be considered carefully, such as vascularization, native cell recruitment, and scar inhibition (Owusu-Akyaw et al., 2019). In the field of IUA treatment, hydrogels are broadly applied as space-filling agents, delivery vehicles for bioactive substances, and 3D structures that organize cells and present stimuli to ensure the repair of the damaged endometrium (Grover et al., 2012). Herein, we summarize the progress in hydrogel research for endometrial regeneration and IUA treatment from the following three aspects; namely, bioactive hydrogels as physical anti-adhesion barriers, in situ drug delivery systems, and 3D cell delivery and culture systems.

Bioactive Hydrogels as Absorbable Physical Anti-Adhesion Barriers in IUA

Generally speaking, the main methods for the treatment of IUA are transabdominal or hysteroscopic resection of the adhesion; while but the former creates appreciable tissue injury, the latter has a high risk of recurrence, and postoperative uterine cavity wounds often quickly form new adhesions (Bhandari et al., 2015). Therefore, positive preventive measures should be taken for IUA after operation. Currently, strategies to reduce the rate of recurrence and improve reproductive outcomes in cases of severe IUA include the use of an IUD, Foley balloon (FB), amnion graft applied over a FB, and an oxidized and regenerated cellulose adhesion barrier (INTERCEED®), with the aim of maintaining the uterine lumen (Ugboaja et al., 2017). However, there are several shortcomings of IUD and FB in preventing the recurrence of IUA. IUD has a potential risk factor of infection (Sun et al., 2018a), and is only effective in preventing re-adhesion (Chi et al., 2018). To solve the problem, some studies investigated the potential benefits of hydrogels as physical anti-adhesion barriers to prevent recurrence in IUA after hysteroscopic adhesiolyis (Table 1).

Compared with patients using an IUD alone, the combination of an oxidized, regenerated cellulose adhesion barrier (INTERCEED®) and an IUD could regain an adhesion-free...
uterine cavity and significantly shorten the conception interval (Cai et al., 2017). A comprehensive treatment strategy of intrauterine injection of HA hydrogel combined with a balloon urinary catheter and IUD insertion, and oral estrogen following hysteroscopic adhesiolyis, was investigated in moderate-severe IUA patients (Xiong et al., 2020). The results indicated that this combined strategy prevented the recurrence of IUA to a certain extent. Moreover, evidence indicated that the combination of IUD and anti-adhesion hydrogel had a more satisfactory effect than applying an IUD alone (Can et al., 2018; Li et al., 2019). Lin et al. (Lin et al., 2013) found that the insertion of an intrauterine balloon or IUD was more effective than the use of HA gel alone in the prevention of IUA reformation. Furthermore, Pabuçcu et al. (Pabuçcu et al., 2019) found that endometrial thickness was significantly enhanced when applying a new crosslinked hyaluronan gel with an IUD after hysteroscopic adhesiolysis. Although many studies have shown that an IUD combined with hydrogels can reduce IUA recurrence, in some cases the clinical pregnancy rate of these patients is not improved. Therefore, in-depth research is needed to explore whether the application of hydrogels can improve pregnancy outcomes for infertile patients. It may be that the pure hydrogels only play an anti-adhesion role, and are unable to promote the regeneration and functional recovery of the endometrium. Therefore, it is necessary to optimize the design to prepare functional bioactive hydrogels that are conducive to endometrial repair and functional regeneration of the endometrium.

Huang et al. (Huang et al., 2020b) tested patented intrauterine scaffolds made of amino acids as a barrier in the treatment of recurrent IUA with poor prognosis. The results showed that the menstrual volume returned to normal, and the recovery rate of amenorrhea increased significantly after scaffold implantation. However, the recurrence rate of IUA was still up to 25%, which indicated that although intrauterine scaffolds may be an effective therapy, they still need to be further optimized.

In addition, although ordinary HA gel can mechanically support the uterine cavity, because of its rapid degradation it cannot stay inside the uterine cavity for a prolonged duration, therefore the anti-adhesion effect is not satisfactory (Azumaguchi et al., 2019). A meta-analysis addressing the use of HA gel to prevent IUA after intrauterine surgery showed that HA gel can reduce the incidence of re-adhesion in mild cases following intrauterine surgery (Fei et al., 2020; Zheng et al., 2020). While there was also research showed that the application of auto-cross-linked HA gel was unable to reduce the recurrence rate of IUA following hysteroscopic adhesiolysis (Zhou et al., 2021). ABT13107, a newly developed thermo-responsive sol-gel made from non-animal derived HA and synthetic poloxamers, had been studied regarding preventing the recurrence of IUA (Lee et al., 2020). Moreover, a randomized double-blind controlled clinical trial (RCT) was conducted to evaluate the efficacy and safety of auto-crosslinked HA gel for preventing IUA after hysteroscopic adhesiolysis; the results indicated that auto-crosslinked HA gel may be able to reduce IUA, decrease adhesion severity, and improve menopause postoperatively (Xiao et al., 2015). A meta-analysis exploring whether the application of HA gel can reduce the recurrence rate of IUA after hysteroscopic adhesiolysis showed that HA gel could reduce the recurrence rate of IUA, but had no significant effect on postoperative pregnancy rate (Fei et al., 2019). In addition, another study compared the effect of adjuvant therapy on the prevention and treatment of IUA and found that an alginate hyaluronate-carboxymethylcellulose membrane (ACH) was most effective in preventing IUA progression (Yan and Xu, 2018). Chenite et al. indicated that sodium hyaluronate or a hydrogel prepared using hydroxybutyl chitosan (HBC) and polyphosphate can reduce the occurrence of postoperative IUA. However, owing to the lack of viscosity of sodium hyaluronate or HBC hydrogel, they are not easily retained in the uterine cavity. Thus, it cannot completely isolate uterine wounds, which affects its ability to prevent IUA. Therefore, after developing and optimizing hydrogels with potential anti-adhesion ability combined with a physical anti-adhesion barrier for IUA, bioactive hydrogels overcome some drawbacks of original natural hydrogels, the prevention and treatment have great clinical significance (Kowalski et al., 2019).

| Physical barrier | Biomaterial | Model | Strength | References |
|-----------------|-------------|-------|----------|------------|
| IUD + Hydrogel  | New crosslinked hyaluronan gel | Human | The application of new crosslinked hyaluronan gel significantly reduces IUA formation | Li et al. (2019) |
| Hydrogel        | New crosslinked hyaluronan gel | Human | New crosslinked hyaluronan gel appears to be able to reduce the formation of IUA | Can et al. (2018) |
| IUD + Hydrogel  | New crosslinked hyaluronan gel | Human | Better endometrial thickness values were observed in those who received new crosslinked hyaluronan gel either alone or in combination with an IUD | Pabuçcu et al. (2019) |
| Hydrogel        | ABT13107    | Human | ABT13107 is non-inferior to the highly viscous hyaluronic acid anti-adhesive barrier in IUA formation after hysteroscopic surgery | Lee et al. (2020) |
| FB + Hydrogel   | Auto-crosslinked hyaluronic acid gel | Human | Auto-crosslinked HA gel could be able to reduce IUA, decrease adhesion severity, and improve menopause postoperatively | Xiao et al. (2015) |
| Hydrogel        | Hyaluronic acid gel | Human | Reduce the risk of postoperative IUA | Tafti et al. (2021) |
(Kwon et al., 2018). The application of bioactive hydrogels in IUA endometrial tissue regeneration is still in the exploratory stage. However, the use of bioactive hydrogels as a local drug delivery system to release specific drugs for regulating the specific pathological environment of IUA has great potential for application in endometrial regeneration. Hydrogels have attracted noticeable interest for their use in drug delivery owing to their unique physical properties. The high porosity that characterizes hydrogels can easily be adjusted by controlling the density of cross-links in their matrix and their affinity to water (Ávila-Salas et al., 2019). Their porous structure also allows drugs to be loaded and then released. The topical application of hydrogels can effectively be used in drug delivery that can help to alleviate the symptoms of many pathological conditions.

**Bioactive Hydrogels as in situ Drug Delivery Systems Through Vaginal Administration**

The large surface area and abundant blood flow in the vagina are excellent for promoting drug absorption (Ci et al., 2017). Vaginal administration has no first-pass effect and has low enzyme activity; therefore, drugs can perform local treatment, and can also enter the systemic circulation. Thermosensitive hydrogels are easily administered vaginally, and are administered in a low viscous form at room temperature (Domiński et al., 2019). After entering the vagina, the hydrogel spreads rapidly into the folded area of the vaginal mucosa and forms a gel at body temperature (Liu et al., 2017). Stromal cell-derived factor-1α (SDF-1α) is a chemokine protein with the ability of recruiting endogenous cells, that can accelerate the regeneration of multiple tissues. In order to obtain a superior effect, Qi et al. (Wenbo et al., 2020) prepared an SDF-1α-loaded-chitosan-heparin hydrogel with controlled drug release ability. Seven days after treatment, uteri treated with SDF-1α incorporating a hydrogel gave both mechanical support to the uterine cavity and also initiated regeneration of the endometrium by recruiting c-kit positive stem cells to the injury site. Cai et al. (Cai et al., 2019) studied the incorporation of SDF-1α within a silk fibroin-bacterial cellulose (SF-BC) membrane that promoted the regeneration
of a full-depth uterine injury and also improved the pregnancy outcome of the damaged uterus.

Aloe has been reported as an ideal organic component to mix with poloxamer to form a more biologically friendly thermosensitive hydrogel system (Baghersad et al., 2018). Aloe-poloxamer (AP) hybrid hydrogel has been fabricated for treating IUA, which can achieve a better therapeutic effect with prolonged retention time within the uterine cavity compared to poloxamer gel. This study showed that AP hydrogel can serve as a biologically active delivery carrier for $\beta$-estradiol and that it can synergistically promote morphological, structural, and functional repair of the injured uterus (Yao et al., 2020) (Figures 1A–D).

Drug Delivery Through Injectable Bioactive Hydrogels

The use of injectable hydrogels, where the solution–gel transition occurs at the site of administration, may shorten the duration of treatment, reduce the risk of infection of the implanted site, and prevent scarring (Asai et al., 2012). To date, various studies have investigated the possibility of this therapy in animal models (Sun et al., 2018b; Jian et al., 2018; Jiang et al., 2019). The injectable SDF-1$\alpha$ release hydrogel exhibited long-term recruiting of hematopoietic stem cells (HSCs) and achieved better effects than a one-off injection of SDF-1$\alpha$ solution. This hydrogel could be a candidate for uterine injury healing and other wound dressing drug delivery systems. Because of their biocompatibility, biodegradability, and tissue adhesion, chitosan and HA are also suitable as a matrix for injectable in
situ hydrogels for drug, gene delivery, and tissue repair through invasive surgery (Lu et al., 2019).

The role of new bioactive hydrogel scaffolds, such as heparin-poloxamer (HP), and gelatin meth acryloyl (GelMA), in uterine cavity repair has attracted growing attention, and they can be used as sustained-release drug delivery systems to effectively promote endometrial repair in IUA animal models (Wu et al., 2019). Zhang et al. (Zhang et al., 2017) injected the combination of different drugs (hormone or growth factor) with thermosensitive, sustained drug delivery systems based on HP hydrogels to rat’s uteri (Figures 2A–D). The study demonstrated that the encapsulation of 17β-estradiol into HP hydrogels can increase the drug concentration at the injured area while reducing the blood concentration, because it can be administered by in situ injection. Keratinocyte growth factor (KGF), a potent repair factor for epithelial tissues, had been encapsulated into HP hydrogels, and the KGF-loaded HP hydrogel delivered a sustained release of long duration and prolonged retention of the encapsulated KGF in the injured endometrium, thus improving endometrial recovery in a rat model compared to the administration of KGF alone (Figures 3A–E) (Xu et al., 2017). However, at present, the above synthetic biological scaffolds are generally expensive and have not been widely used in clinical settings. In animal experiments, most injectable bioactive hydrogels are administrated through the myometrium rather than the cervical canal. However, gynecologists usually apply anti-adhesion hydrogel through the cervical canal for IUA patients after hysteroscopic adhesiolysis. Though these hydrogels are all injectable, different administrations make it is not clear whether these patients can benefit from injectable bioactive hydrogels.

Bioactive Hydrogels as 3D Cell Delivery and Culture Systems in IUA

Stem cell therapies have been demonstrated to be promising and multifunctional alternatives to traditional drug therapy. To date, numerous maladies have been modeled in animals to test the efficacy and safety of a range of cellular therapies. However, the difficulty lies in the normal expansion of stem cells in cell therapy. There are two crucial factors that influence stem cell expansion, which are biochemical composition and the physical properties of the matrix (Madl et al., 2018). One crucial physical regulator of stem cell fate is matrix stiffness. Interestingly, bioactive hydrogels can be fabricated as culture systems with a tunable stiffness that encompasses a physiological range. Owing to their high-water content, porosity, and soft consistency, bioactive hydrogels can closely simulate natural living tissue. In addition, by adjusting the biochemical composition of the hydrogel or encapsulated drugs, they can
Table 2 | Summary of bioactive hydrogels in cell therapy in IUA.

| Cell                        | Biomaterial                        | Model              | Molecule/pathway                                      | Strength                                                                 | References               |
|-----------------------------|------------------------------------|--------------------|-------------------------------------------------------|---------------------------------------------------------------------------|--------------------------|
| AdMSC-derived exosomes      | Hyaluronic acid fibrinogen/thrombin hydroygel (HA + F + T50 gel) | Sprague-Dawley rats | Increased expression of PECAM and IGF-1                 | VEGF and LIF expression are also increased                               | Xiao et al. (2018)       |
| MSC-Sec                     | Crosslinked hyaluronic acid gel    | Sprague-Dawley rats | -                                                     | Creates a sustained release system                                        | Liu et al. (2019)        |
| BMSCs                       | Hyaluronic acid fibrinogen/thrombin hydroygel (HA + F + T50 gel) | Sprague-Dawley rats | Increased expression of HuNu and vimentin in a IUA-induced rat model | Effective in preventing fibrosis with improved regeneration of endometrium and endometrial glands in the rat model of IUA | Chen et al. (2019)       |
| Oral mucosal epithelial cells | Lyophilized amniotic membrane     | Sprague-Dawley rats | VEGF expression is increased                          | Increased number of endometrial glands and reduced area of fibrosis     | Liu et al. (2020a)       |
| hUCMSC                      | Collagen                           | Sprague-Dawley rats | Increased expression of HuNu and vimentin in a IUA-induced rat model | Increased number of endometrial glands and reduced area of fibrosis     | Liu et al. (2020a)       |
| UC-MSCs                     | Collagen                           | Human              | -                                                     | 38.4% participants had a successful pregnancy                             | Cao et al. (2018)        |
| UC-MSC-derived exosomes     | Collagen                           | Sprague-Dawley rats | Increased expression of the estrogen receptor α/progesterone receptor | Endometrium regeneration                                               | Xin et al. (2020)        |
| hMSCs                       | Alginate and gelatin              | Sprague-Dawley rats | Integrin αβ3 and LIF were significantly higher in the cell-loaded scaffold group | Promoted the recovery of the endometrial histomorphology (endometrial tissue and gland regeneration) and the regeneration of endometrial cells (stromal cells and epithelial cells) and endometrial cells | Ji et al. (2020)         |
| ADSCs-derived exosomes      | PEG                                | Sprague-Dawley rats | Significant increases in VEGF, LIF, αvβ3, and IGF-1 expression | Promote HUVEC proliferation, migration, and tube formation in vitro | Pan et al. (2020)        |

hAECs: human amnion epithelial cells; bFGF: basic fibroblast growth factor; VEGF: vascular endothelial growth factor; IGF-1: insulin-like growth factor-1; COL1A1: collagen type I alpha 1; TIMP-1: tissue inhibitor of metalloproteinase-1; TGF-β: transforming growth factor-β; PDGF-C: platelet-derived growth factor-C; THBS1: thrombospondin-1; CTGF: connective tissue growth factor; ER-β: estrogen receptor β; 2%AluECMNPs/AP hydrogel: nanoparticulate decellularized uterus (ECMNPs) encapsulated β-estradiol thermosensitive aloe-poloxamer hydrogel; MenSCs: menstrual blood stem cells; BMSCs: bone marrow-derived stem cells; dEMSCs: decidualized endometrial stromal cells; MSC-Sec: mesenchymal stem cell-secretome; UC-MSCs: umbilical cord-derived mesenchymal stromal cells; ADSCs: adipose stem cells.

also regulate the fate of stem cells. Therefore, functional bioactive hydrogels make it easier to transport stem cells to injured the endometrium with good vitality and differentiation potential. At present, there are ongoing studies on the advantages and disadvantages of different bioactive hydrogels applied in stem cell therapies in IUA. A summary of bioactive hydrogels in stem cell therapy in IUA is listed. The advantages of the mentioned hydrogels are also listed (Table 2). According to Table 2, the endometrium regeneration associated molecules are well improved. Another highlight of bioactive hydrogel used in stem cell therapy is that it can decrease uterine fibrosis rate and promote endometrial glands growth.
Hyaluronic Acid

HA may be an appropriate hydrogel for endometrial regeneration therapeutics. Nevertheless, HA gels have a natural half-life of just 1–2 days (Carruthers and Carruthers, 2007). Moreover, the half-life may be even shorter in the uterus owing to aqueous dilution. In order to optimize the short half-life of natural HA, Kim et al. (Kim et al., 2019) constructed endometrium-tailored HA/fibrin composite hydrogel and demonstrated the significant regenerative effects of the in vitro processed isotopic cells encapsulated in composite scaffold materials. The expression of PECAM and IGF-1, which are neovascularization- and decidua-specific genes, were significantly increased. Vascular endothelial growth factor (VEGF) and leukemia inhibitory factor (LIF) expression were also increased in the HA-fibrin-thrombin 50 mg (HA-F-T50) composite hydrogel. Moreover, an attempt at pregnancy was possible as early as 2 weeks after treatment, and successful in vivo embryonic implantation and development in the regenerated recipient model was confirmed at 7 days after embryo transformation in a murine uterine synchiea model. Despite limited evidence of its efficiency in the human body, further studies should be carried out to optimize this promising technology.

Liu et al. (Liu et al., 2019) fabricated a control-released, intrauterine-administered mesenchymal stem cell-secretome (MSC-Sec)-loaded, crosslinked HA gel, that was developed as a therapy to restore injured endometrial morphology and fertility in a rat model of AS (Figures 4A–D). This crosslinked HA gel can be used in stem cell therapy because it can bypass the tumorigenic risks associated with stem cell therapies and help maintain the promising effect of MSC-Sec on endometrial and endothelial cells.

Other Hydrogels

Apart from HA hydrogel, Pluronic F-127 is a synthetic Food and Drug Administration (FDA)-approved compound which has several advantages, including low toxicity, biocompatibility, and thermo-reversibility. It is widely used in drug delivery and in vivo tissue engineering because it can form into a hydrogel at physiological temperatures. BMSCs and Vitamin C encapsulated Pluronic F-127 hydrogel was prepared to promote restoration of damaged IUA endometrium in vivo; the vitamin C promoted BMSCs survival and growth in the hydrogel (Yang et al., 2017).

In another study, Liu et al. (Liu et al., 2020b) used collagen scaffold loading with human umbilical cord mesenchymal stem
cells (hUCMSCs) to increase the number of endometrial glands and reduced the area of fibrosis, suggesting that the combination of the collagen scaffold and hUCMSCs may be an alternative approach for treating IUA. Moreover, the hUCMSCs-collagen scaffold had passed through a phase I clinical trial and obtained a satisfactory outcome, where around 38.4% of participants underwent a successful pregnancy without complications associated with stem cell therapy (Cao et al., 2018).

In terms of 3D bioactive hydrogel scaffolds, chitosan is a biopolymer with a unique set of biological and physicochemical properties (Pan et al., 2020). Thus, Ksenia et al. (Carruthers and Carruthers, 2007) successfully formed different scale structures of chitosan and further tested their biomechanical quality and biocompatibility to demonstrate the prospects of their broad biological application (Figures 5A–D). Furthermore, Ji et al. (Ji et al., 2020) studied the application of 3D bioprinting of a human iPSC-derived MSC-loaded scaffold for repair of the uterine endometrium. The results showed that the endothelial cells and endometrial cells regenerated, and the dysfunctional endometrium was partially restored (Figures 6A–E).

Recently, bioactive hydrogels laden with stem cell exosomes have been developed for endometrial regeneration. The advantages of hydrogel treatment include the potential for sustained exosome release, resulting in the maintenance of higher local concentrations of pharmacologically important compounds for extended periods, and reducing the need for repeated dosing in a clinical setting, thus making them a promising candidate for use in exosome-based endometrial repair applications (Pan et al., 2020). Lin et al. (Lin et al.,...
developed a microenvironment-protected exosome-hydrogel for endometrial regeneration (Figures 7A–C). This PEG hydrogel was injectable, antibacterial, and facilitated controlled ADSC-exo release in order to promote endometrial regeneration.

CURRENT LIMITATIONS AND FUTURE PERSPECTIVES

Despite their many advantageous properties, hydrogels also have several limitations. With the continuous progress of related research, the disadvantages of natural hydrogels are being gradually revealed. Although some natural hydrogels such as HA and alginate have already shown superiority over other hydrogels, the opportunity to improve current hydrogel implementations is still very promising. Researchers are gradually improving the structure of hydrogels by changing the polymer type and optimizing fabrication methods. The low tensile strength of many hydrogels limits their use in load-bearing applications, resulting in displacement of the hydrogel from a targeted local site. Because the hydrophobic compounds have limited loading quantity and homogeneity in hydrogel matrices, hydrogels are limited to carrying hydrophilic drugs (Zagórska-Dziok and Sobczak, 2020).

Physical Anti-Adhesion Barriers

According to the published research (Du et al., 2020), anti-adhesion hydrogels are usually placed in the uterine cavity to prevent recurrence after operation. However, the effect of ant-
adhesion hydrogels alone is less satisfactory than that of a combination of hydrogels and IUD. The reason why hydrogels cannot become the first-line therapy in endometrial regeneration is that most hydrogel applications in the prevention of IUA are still less satisfactory than a traditional IUD. Moreover, hydrogels do not have sufficient mechanical strength to shape a normal uterine cavity throughout a complete treatment. Unlike traditional IUD, some hydrogels in liquid are able to transform to sodium gel after entering uteri, which may help more efficient formation of the uterine cavity in various sizes. It is known that most IUA patients in childbearing age hope to improve fertility through IUA treatment. However, some bioactive hydrogels as physical barriers cannot assist in the formation of a normal functional endometrium for embryo implantation because they cannot meet the requirement of excellent cell culture system and sufficient mechanical strength. Therefore, future research needs to address the optimization of bioactive hydrogels as physical barriers to increase their strength in shaping a normal uterine cavity while still maintaining the capacity of controllable degradation.

**Drug Delivery Systems**

As mentioned previously, hydrogels can be applied as in situ drug delivery systems for sustained release drugs to promote endometrial regeneration in patients with IUA. Compared with traditional hormone replacement therapy, this method may reduce the incidence of related risks like breast hyperplasia caused by oral estrogen in treating IUA. Several experiments that demonstrate the effectiveness of a hydrogel-based drug delivery model in treating IUA have used animal models, and there has been limited evidence of clinical practice in human (Wang et al., 2020b). Whether hydrogels as in situ drug delivery systems are more beneficial than traditional drug delivery is still not clear in humans. Moreover, the hydrated nature of hydrogels can make terminal sterilization difficult and time-consuming (Lima et al., 2020). In the future, hydrogels with multiple concurrent functions may be a satisfactory way of treating IUA, which can provide both mechanical structure in forming the uterine cavity, and also be a stable in situ drug delivery system.
Stem Cell Therapy

Because of their structural features, bioactive hydrogels are widely used as 3D cell culture systems. With the recent broader understanding that stem cells and stem cell derivatives can promote endometrial repair, the value of 3D hydrogel scaffolds that can simulate the living environment of cells and promote the transmission of stem cells and their secretions has become particularly prominent. Therefore, some studies have explored the application of hydrogels in IUA cell therapy. Many animal experiments have shown that, compared with traditional cell therapy, the application of hydrogels as a 3D culture environment produces superior outcomes in endometrial regeneration, and the improvement of endometrial receptivity is evident. However, despite exhibiting promising results in animal experiments, the application of hydrogels in stem cell therapy is still largely restricted to the experimental stage owing to its critical pitfalls and drawbacks such as safety issues, poor cell survival, and high cost.

Currently, one of the biggest concerns regarding stem cell therapy is the tumorigenic complications caused by uncontrollable stem cell implantation. Thus, the future study of hydrogels used in stem cell therapy could focus on controllable cell growth offering a better therapeutic approach while avoiding related complications. Moreover, the existing cell loaded hydrogels face the problem that the supply of oxygen and nutrients is limited by diffusion kinetics. For example, 3D printed hydrogels still require surface modification, such as chemical methods to carry integrin (for endometrial cells) and increase the survival rate of the seeded cells, by inducing cell-oriented differentiation, or by maturing the scaffold in a bioreactor before implantation (Wen et al., 2019). Therefore, selecting and optimizing bioactive hydrogels is important in obtaining better results and for their application in treating IUA.

CONCLUSION

IUA is a common endometrial disease and one of the main causes of infertility in women of childbearing age. Owing to their excellent properties—such as good biocompatibility, degradability, and controlled drug release—bioactive hydrogels play an important role in the prevention and treatment of IUA and have great potential for application in the clinical setting. Bioactive hydrogels can be used as physical anti-adhesion barriers, and can also act as drug delivery systems for hormone drugs, multiple factors, and 3D cell delivery and culture systems in IUA treatment. These characteristics show that a combination of hydrogels and traditional intrauterine adhesion treatment can have a significant therapeutic effect on IUA and improve pregnancy success rates. While most researches ongoing are based on animal experiments, the mechanism behind the therapeutic effect of bioactive hydrogel in IUA treatment is not clear. The optimization on the safety and effectiveness of bioactive hydrogels is valuable to researchers in the future.

AUTHOR CONTRIBUTIONS

JW and YL conceptualized, designed, and wrote the manuscript. CY and YX drafted the manuscript, reviewed the literature. TJ and XC designed the figure and table. YL, JT, and SH revised the manuscript. All authors approved the final version of the manuscript.

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