Endometriosis: Survey of Current Diagnostic and Therapeutic Options and Latest Research Work

I. Juhasz-Böss1, M. W. Laschke2, F. Müller1, P. Rosenbaum1, S. Baum1, E. F. Solomon1, U. Ulrich3

Abstract

Endometriosis is one of the most frequent benign diseases in women of child-bearing age. The main symptoms are chronic upper abdominal pain and infertility. However, the aetiology and pathogenesis of endometriosis are as yet insufficiently clarified. Thus, therapy is mainly symptomatic with laparoscopic surgery being the gold standard. The aim of drug therapy is to achieve a hypo-oestrogenic condition. In cases of severe endometriosis and a desire to have children there is often an indication for assisted reproduction. The present article illustrates almost all current aspects on the diagnosis of and therapy of endometriosis. From the clinical viewpoint, emphasis is placed on the rare cases of deeply infiltrating endometriosis that are, however, accompanied with a high morbidity. Current therapeutic options in cases of infertility are also presented in more detail. Furthermore, special attention is paid to the recent research results from both clinical and basic research fields in order to demonstrate our current knowledge on the pathogenesis and, where possible, potentially related therapeutic options.

Introduction, Definitions and Epidemiology

Endometriosis is one of the most frequent benign diseases of the fertile woman and is defined as the presence of endometrium outside of the uterine cavity. The prevalence rates given in the literature vary widely. It is estimated that in Germany about 40,000 new cases occur each year [1]. Risk factors for endometriosis are, among others, the duration of period bleeding, length of the menstrual cycle, number of pregnancies, number of miscarriages and smoking status [2]. The pathogenesis of the disease is even today not completely clarified. Although endometriosis is a benign disease, it can spread to other organs through infiltrative growth and thus require extensive surgery. The main symptoms of the disease are sterility and chronic pain [3,4].

Current therapeutic concepts are mainly aimed at suppression of ovarian function and analgesia in addition to the operative management. For this purpose above all gestagen-rich contraceptives, pure gestagen and GnRH analogues are employed. For pain therapy NSAR are used preferentially. In spite of the use of known therapeutic

Zusammenfassung

strategies in some cases insufficient relief of the endometriosis-related complaints or disease recurrence can occur.

The disease is thus also of economic relevance. Beside the medical expenses, above all the patients’ inability to work is appreciable [1,5]. Repeated surgery, the in part chronic course as well as the delayed diagnostic confirmation after year-long symptoms and complaints results in a high medical expenditure [6].

The present article aims to give an actual survey of the diagnostic and therapeutic procedures as well as to present the latest research results on the subject of endometriosis.

Aetiology of Endometriosis: Classic Theories and Latest Research Approaches

Although endometriosis is one of the most investigated gynaecological clinical entities, many aspects of the aetiology and pathogenesis of the disease have still not been completely clarified and thus represent a focus of current research on endometriosis. The classical theories on the development of endometriosis include the “coelom metaplasia theory” (according to Meyer) and the “theory of embryonic cell remains” according to which the endometriosis lesions develop from tissue of embryonic origin, as well as the “theory of lymphovascular metastasis”, which proposes that endometrial cells, similar to tumour cells, spread via lymphatic pathways or blood vessels and that in this way endometriosis lesions can develop in various positions in the body [7]. Today the most widely accepted theory on the development of endometriosis is, however, the “implantation theory” according to Sampson [8]. This is based on the assumption that during menstruation, vital endometrium reaches the abdominal cavity from the uterus in a retrograde manner via the tubes. Favourable conditions there lead to implantation of tissue fragments on the peritoneum and thus to the development of endometriosis. However, retrograde menstruations are certainly physiological and can be detected in up to 90% of all women who undergo laparoscopy [9]. Thus there is no doubt that the development of endometriosis depends on many other pathogenic mechanisms, thus rendering it a typical multifactorial clinical entity. Numerous studies have in the meantime indicated that there is a genetic predisposition for endometriosis [10]. For this reason genome-wide association studies are underway in order to identify the corresponding genetic factors. Although in the course of these studies 3 potential gene loci have already been found that are with the highest probability associated with the occurrence of endometriosis in Japanese and European women, it is still necessary to confirm such results in a larger number of cases and to correlate the obtained data with the various clinical subtypes of endometriosis [11]. In addition epigenetic studies are needed to clarify how certain environmental factors influence the disease. In this context, dioxins must be considered as potential disease-triggering environmental toxins [12]. Furthermore, certain components of foodstuffs such as, e.g., fatty acids and their derivatives, are being discussed as factors influencing the disease [13]. Various immunological aspects also play an important role in the pathogenesis of endometriosis, whereby the disease shows marked similarities with typical autoimmune diseases [14]. Thus, it was shown that in the course of endometriosis autoantibodies against endometrial antigens such as, e.g., transferrin or alpha 2-HS glycoprotein, are formed, and which can often in part be responsible for the infertility frequently observed in endometriosis patients [15,16]. Also concentration fluctuations of cytotoxic and activated lymphocytes peripheral blood during the menstrual cycle appear to play a certain role. The fluctuations of regulatory T cells detected during endometriosis can be attributed to a changed immune response [17]. Furthermore, endometriosis is characterised by a chronic inflammatory reaction with elevated concentrations of inflammatory cytokines in serum and the peritoneal fluid [18]. Elevated numbers of macrophages, dendritic cells and natural killer cells are seen in the peritoneum which, as for as yet insufficiently investigated reasons, are however not able to recognise and degrade the endometrium tissue scattered in the abdominal cavity [19]. This chronic inflammatory reaction, in turn, leads to an increased production and an inhibited degradation of reactive oxygen species [20]. Accordingly, the administration of substances with an antioxidative action could represent a meaningful supplement to the current therapeutic options.

The basal parts of the endometrium contain endometrial stem cells and progenitor cells [21], which explains the high regeneration potential of this tissue. This is at present under discussion as a further cause for the occurrence of endometrium lesions. Thus, it is assumed that stem cells reach the abdominal cavity via retrograde menstruation and under suitable conditions can then differentiate into endometriosis lesions [22]. This assumption is supported by the observation that the glandular cells of some endometriosis lesions are of monoclonal origin [22]. In addition, it is possible that also extraterine, circulating stem and progenitor cells from bone marrow contribute appreciably to the formation of endometriosis lesions. Thus, it has been shown in a mouse model that not only glandular but also stromal cells from bone marrow cells can develop into endometriosis lesions [23]. Furthermore, up to 37% of all endothelial cells in endometriosis lesions are derived from circulating endothelial progenitor cells [24]. A major prerequisite for the long-term survival of endometriosis lesions is angiogenesis, i.e., the generation of new blood vessels [25]. Only in this way can the scattered endometrium tissue, which is initially ischaemic outside the uterus, be adequately supplied with oxygen and nutrients. For this reason, especially the early stages of endometriosis lesions exhibit a reddish colouration caused by a high density of blood vessels and vessel dilatations [26,27]. Furthermore, in the past few years numerous angiogenic growth factors have been identified – above all “vascular endothelial growth factor” (VEGF) – which is produced and released in increased amounts in endometriosis, and regulates the ingrowth of new blood vessels in endometriosis lesions [28]. Interestingly, endometrium in the uterus of endometriosis patients already exhibits an elevated angiogenic activity which can favour the process of creation of new endometriosis lesions [29]. Endometriosis thus belongs to the group of angiogenic diseases along with rheumatoid arthritis, psoriasis, diabetic retinopathy as well as tumour growth and metastasis [29]. Accordingly in recent years increasing attention has been paid to analysing the control mechanisms that govern the formation of new blood vessels in endometriosis lesions, in order to identify new targets for a targeted, anti-angiogenic endometriosis therapy [30,31]. In addition, experimental in-vitro and in-vivo studies have already detected numerous members of various substance classes that exert an anti-angiogenic effect on endometriosis lesions. These include growth factor inhibitors, endogenous angiogenesis inhibitors, fu-magillin analogues, statins, cyclooxygenase 2 inhibitors, phytotherapeutic agents, immune modulators as well as dopamine agonists [32]. Further clinical trials are needed to clarify to what extent these active principles can be used for an anti-angiogenic
endometriosis therapy without causing severe side effects in the afflicted patients.

Another area of current research focusses on the mechanisms that contribute to the pain symptomatic of endometriosis. Major contributing factors are considered to be long-term cyclic bleeding of the oestrogen-dependent endometriosis lesions with consecutive inflammatory reactions as well as irritation and invasion of pelvic nerves [33]. Recent investigations, however, have shown that pain transmitting nerve fibres can grow together into endometriosis lesions [34], this is also designated as “neuroangiogenesis” [35]. These nerve fibres in turn increase the pain sensation in the central nervous system [35]. If it were possible – as has already been demonstrated in animal experiments [36] – to inhibit this process by targeted drug measures, we should be able in the future to develop appreciably more effective pain therapies having considerably fewer side effects than the conventional hormone treatment options.

Summarising the current research results, we see that endometriosis is a complex, multifactorial disease for which the aetiology and pathogenesis have not yet been completely clarified. The research findings of the past 10 years, however, do show that molecular and cellular pathomechanisms which contribute to the development of the disease are increasingly being identified. This should provide the possibility in future to develop better targeted, new preventive and therapeutic treatment strategies.

**Diagnostic, Clinical Picture and Therapy for Endometriosis**

As recommended in the current guidelines and for the sake of better clarity, the differing manifestations of endometriosis are discussed separately although they often occur in combination [1]. First of all, the following aspects are valid for all manifestations:

- **Laparoscopy:** Laparoscopy (LSC) is a central component in the diagnosis of and therapy for endometriosis. Indication for LSC is: pain, organ changes and/or sterility. Surgical removal of the endometriosis lesions is considered to be the gold standard, on the one hand to control the symptoms and on the other hand for histological work-up [37]. It should be noted that asymptomatic endometriosis in a patient not desiring to have children does not require any surgical or drug treatment [1]. Examination of the case history and the additional use of psychological questionnaires provide valuable information prior to the diagnostic/curative operation and can provide indications for the presence of an endometrial disease and thus support an optimal counselling for the symptomatic patient in the phase of decision-making for surgical clarification [4].

- **Localisation:** Endometriosis preferentially attacks the lesser pelvis and here above all (in order of decreasing frequency) the pelvic peritoneum, the ovaries, the sacrouterine ligament, the rectovaginal septum as well as extragenital manifestations such as, e.g., the rectosigmoid or urinary bladder. Extraperitoneal manifestations are seldom [1].

- **Staging:** In clinical routine endometriosis is subdivided according to its localisation into endometriosis genitalis externa and interna as well as extragenitalis [1]. Unfortunately endometriosis is not uniformly classified, furthermore the current staging is not satisfactory. The most widely used and most frequently used in reproduction medicine is the rASRM classification of the American Society for Reproductive Medicine (ASRM) [38] (Table 1).

<table>
<thead>
<tr>
<th>rASRM classification of endometriosis (modified from [38]).</th>
<th>stage I (minimal): 1–5; stage II (mild): 6–15; stage III (moderate): 16–40; stage IV (pronounced): &gt;40.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Endometriosis</strong></td>
<td></td>
</tr>
<tr>
<td>Peritoneum</td>
<td></td>
</tr>
<tr>
<td>superficial</td>
<td>&lt; 1 cm</td>
</tr>
<tr>
<td>deep</td>
<td>1–3 cm</td>
</tr>
<tr>
<td>deep</td>
<td>&gt; 3 cm</td>
</tr>
<tr>
<td>Ovaries</td>
<td></td>
</tr>
<tr>
<td>R superficial</td>
<td>&lt; 1 cm</td>
</tr>
<tr>
<td>R deep</td>
<td>1–3 cm</td>
</tr>
<tr>
<td>L superficial</td>
<td>&gt; 3 cm</td>
</tr>
<tr>
<td>L deep</td>
<td></td>
</tr>
<tr>
<td>Adhesions</td>
<td></td>
</tr>
<tr>
<td>partial</td>
<td>&lt; ½</td>
</tr>
<tr>
<td>complete</td>
<td>&gt; ½</td>
</tr>
<tr>
<td><strong>Ovaries</strong></td>
<td></td>
</tr>
<tr>
<td>R thin</td>
<td>&lt; ½</td>
</tr>
<tr>
<td>R dense</td>
<td>&gt; ½</td>
</tr>
<tr>
<td>L thin</td>
<td>&lt; ½</td>
</tr>
<tr>
<td>L dense</td>
<td>&gt; ½</td>
</tr>
<tr>
<td><strong>Tubes</strong></td>
<td></td>
</tr>
<tr>
<td>R thin</td>
<td>&lt; ½</td>
</tr>
<tr>
<td>R dense</td>
<td>&gt; ½</td>
</tr>
<tr>
<td>L thin</td>
<td>&lt; ½</td>
</tr>
<tr>
<td>L dense</td>
<td>&gt; ½</td>
</tr>
</tbody>
</table>

* in cases of complete attack or, respectively, complete occlusion of the fimbria ends: 16

Retroperitoneal and deep infiltrating endometriosis are best described by means of the ENZIAN classification [39,40] (Fig. 1). However, the stage does not correlate with the degree of complaints, some afflicted women are even asymptomatic.

**Tumour marker CA-125:** CA-125 is generally elevated in endometriosis patients. However, this tumour marker has no differential diagnostic relevance. In clinical routine its determination is not recommended either for diagnosis or for course control [1].

**Peritoneal endometriosis**

**Morphology and symptoms**

In cases of peritoneal endometriosis a distinction is made between pigmented (= typical) and not pigmented (= atypical) lesions as well as between red, white and black lesions [26,41] (Fig. 2). These lesions differ above all with regard to their activity, fibrosis, age, etc. It is not clear to what extent the different forms correlate with specific symptoms of endometriosis [42]. Patients with preoperatively pronounced complaints have a higher risk of recurrence than patients with lower pain sensations [43].

**Diagnostics**

Peritoneal endometriosis cannot be detected by sonography. Standard procedure in the field of diagnosis of peritoneal endometriosis is laparoscopy [1]. Histological confirmation should also be attempted for peritoneal endometriosis as mentioned above [1].

**Surgical and drug therapies**

The main objective is the laparoscopic removal of all peritoneal lesions by means of coagulation, vapourisation or excision [1]. It has not yet been clarified as yet which procedure is the most suitable. In answer to this question there is only one randomised controlled trial and the Cochrane analysis published in 2014 [44,45]. The aim of drug therapy is to achieve a hypo-oestrogenic status. Endometriosis implants can be regressively changed by means of...
suppression of ovarian function. The drugs predominately used today are pure gestagens, gestagen-rich contraceptives, GnRH analogues and danazol. GnRH analogues are in such cases more effective than oral contraceptives or gestagens. A reduction of endometriosis-associated complaints can be achieved with all the mentioned substance classes, while GnRH analogues proved to be more effective for dysmenorrhoea and dyspareunia in some trials. Some of these drugs are associated with appreciable and, above all, different side effects. GnRHa should only be administered together with the corresponding protective accompanying drugs ("add-back") because of the consequences of a possible oestrogen deficiency [1].

With regard to the response to hormone therapy it seems that peritoneal endometriosis differs from ovarian and deep infiltrating endometriosis. Duration of therapy with GnRH analogues amounts to 3–6 months. Although a 3-month therapy is equally effective the subsequent recurrence-free interval is then shorter [46]. Recent data from prospective studies have shown that dienogest as maintenance therapy after administration of GnRH can uphold the GnRH-induced effect for at least 12 months [47]. In 2 further RCCCTs equieffective actions of dienogest vs. GnRH analogues for endometriosis-associated pain were observed whereby with regard to clinical tolerance dienogest was better tolerated by the patients [48, 49]. Although frequently used in clinical routine there is no evidence for a positive effect of NSAR on endometriosis-specific complaints [50].

**Ovarian endometriosis**

**Diagnostics**

The ovaries are attacked in up to 50% of all endometriosis patients [1, 51]. Preoperative clarification is based on the clinical ex-
Therapy is adapted to the patient’s family planning wishes. When family planning has been completed hysterectomy (HE) represents the most effective option. Here all common modalities of HE can be applied, but it must be taken into account that a possibly coexisting intraperitoneal endometriosis can usually not be managed by a purely vaginal procedure. A supracervical HE may also be envisioned but should not be performed in case of a concurrent coexisting rectovaginal endometriosis. When planning an HE the operative procedure should be decided upon between surgeon and patient on a case by case basis [1]. For patients still desiring to have children or with a wish to retain their organs, the benefit of an operation has not been demonstrated. Interventional radiological procedures (e.g., embolisation) or MRI-guided focussed ultrasound ablation are still in the experimental stage and should only be applied with the framework of clinical trials [1]. As alternatives to HE, gestagens, hormonal contraceptives and intrauterine, local gestagen-releasing systems may be considered. The therapeutic effect is based on the induction of amenorrhoea. Contraceptives (monophasic formulations) and gestagens should therefore be taken continuously [1].

Deeply Infiltrating Endometriosis

In comparison to peritoneal or ovarian endometriosis, still very little is known about deeply infiltrating endometriosis (DIE) although it is a clinically no lesser relevant form of the disease. In this form it is seen that endometriosis – similar to malignant diseases – can spread beyond organ borders and infiltrate various structures. The thus resulting space-filling lesions and in part invasive growth can lead to indurations and finally to the destruction of functional mobile layers (e.g., between rectum and vagina) through to the occurrence of larger conglomerate tumours and in some cases even to disabling complaints. The fact that, in Germany, statistically an interval of 6 years must pass from the first manifestation of symptoms to the diagnosis of endometriosis is disturbing and may in part be due to frequent misinterpretations and a lack of experience with the deeply infiltrating variant of the disease (see Figs. 3 and 4) [1]. The mostly retroperitoneal growth of DIE is often not directly visible with laparoscopy – due either to the retroperitoneal position or complete Douglas obliteration – and thus it can hardly be defined in the common stage classifications (AFS/rASRM) [57]. DIE should thus additionally be described using the ENZIAN classification (see Figs. 4 and 5) [40]. Frequently affected are the sacrouterine ligament, the parametria, the rectovaginal septum/fornix of the vagina, the rectosigmoid, the appendix, the urinary bladder and sometimes the ureter or also the small intestine.

Symptoms

The symptoms of DIE depend on the afflicted organs or, respectively, structures and vary between no complaints and disability and invalidisation due to severe pain. On the whole and as for endometriosis in general, there is no correlation between the extent of the disease and the severity of the symptom spectrum. A small node in the rectovaginal septum can, for example, make intercourse so painful that it is no longer possible and turn defaecation into a nightmare for the patients, whereas some other patients with stenosed rectum and/or space filling tumours through to frozen pelvis experience no complaints at all. In general, the symptoms of DIE, beside the classical dysmenorrhoea, do of course depend on the pattern of attack. Infiltration of the rectovaginal septum/fornix of the vagina or, respectively, the rectum often leads to dyspareunia and dyschezia, sometimes

Adenomyosis uteri

Diagnostics

In the diagnostic work-up besides gynaecological examinations the transvaginal sonography (TVS) above all and if necessary MRI are most suitable in such cases. TVS is of greater relevance in daily practice [56]. The sonographic criteria for making a diagnosis are poorly delineated heterogeneous areas, in part, cystic intramural alterations, areas of changing echogenicity, irregular halo effects as well as a discrepancy between the anterior and posterior wall findings. Histological confirmation of adenomyosis is in most cases, however, only possible on hysterectomy sample.
also to flatulence and tenesmus and can be accompanied by rectal passage of mucous or blood as well as changes in stool behaviour through to paradox diarrhoea. DIE of the bladder or ureter can – beside hydronephrosis – cause dys- and haematuria. The lack of symptoms never excludes attack of the mentioned organs.

Diagnostics
The suspicion of DIE is based first of all on case history plus the clinical-gynaecological examination, where importance must be placed of an adequate deconvolution of the posterior fornix of the vagina in the speculum settings (see Fig. 3) and a rectal or rectovaginal palpation. Typically among the findings to be recorded – apart from pain trigger points – are visible endometriosis in the vagina, the rough nodular induration of the rectovaginal septum or the parametria/sacrouterine ligaments and, in some cases, urinary retention on sonography. With sufficient experience an infiltration of the rectovaginal septum and the rectum can also be demonstrated by transvaginal sonography [51,56]. The sonographic detection of bilateral endometrioma (especially in the sense of kissing ovaries) can be suggestive of DIE [58].

In the literature magnetic resonance imaging exhibits a high sensitivity for the diagnosis of FDI but is certainly dependent on the investigator’s experience or, respectively, knowledge of the disease but only rarely does it lead to a change of the therapy [59, 60]. In cases of suspected intestinal involvement by DIE colonoscopy is rarely useful since the endometrial infiltration usually stops at the mucosa so that the exclusion of intestinal attack is not possible by rectoscopy. In such cases rectal endosonography is to be preferred [1,59]. Coloscopy must only be used when it is necessary to exclude other differential diagnoses (e.g. colon cancer, diverticulitis, chronic inflammatory bowel disease). DIE of the bladder can often be well visualised by sonography when the bladder is sufficiently full. In such cases, however, cystoscopy prior to surgery is reasonable in order to estimate the distance to the trigonum vesicae and the ureter orifices and, if necessary, to carry out thereby a splinting with a double J catheter.

Therapy
The therapy of choice for symptomatic DIE is resection with healthy margins [61–64]. The operations often include rectum resection (mostly en-bloc with the afflicted rectovaginal septum and the vagina), and partial resection of the sacrouterine ligament and parametria as well as parts of the bladder (see Figs. 4 to 7). Partial ureter resections with new implantation (e.g., psoas-hitch plasty) are much more rarely needed. Hysterectomy is not obligatory and is not wanted by the mostly young patients. The interventions can mostly be performed as laparoscopic or, if necessary vaginal-assisted procedures. Good cooperation between the various disciplines (gynaecology, surgery, urology) is a necessity and accordingly is required for the certification of specialised surgical centres by the Stiftung Endometrioseforschung, the Europäische Endometrioseliga and the Endometriose-Vereinigung-Deutschland e.V.
Urinary retention – due to extrinsic or intrinsic DIE of the ureter – is the only acceptable absolute indication for surgery in the German-language guidelines since it is essential to prevent persisting damage to the kidneys [1]. An intestinal perforation or ileus due to stenosing DIE of the bowels can give rise to an urgent indication in a similar manner [65].

In the case of DIE diagnosed near to the ureter or its recent operative management long-term biannual sonography of the kidneys is recommended in order to recognise the potentially slowly but then also often asymptomatic development of urinary retention in a timely manner [1].

The decision for or against operative removal of DIE is – when as in most cases no absolute indication is given – made together with the patient on the basis of the symptoms and the level of suffering. In most cases an estimation of the extent by means of an at first purely diagnostic laparoscopy is helpful for both physician and patient. In no case may the decision be made without sufficient counselling and time for consideration since extensive rehabilitating resection with participation of the bowels and/or bladder interventions do not guarantee either therapeutic success or freedom from de novo complaints resulting from surgery (late postoperative problems due to disorders of defaecation or to scar and adhesion complaints) [62,66]. An appropriately high burden of suffering is a decisive prerequisite for rehabilitating operations.

Critical in this context is always the wish for removal of an asymptomatic DIE merely on the basis of sterility. In the literature higher rates of not only spontaneous but also of IVF-induced pregnancies after complete resections in such cases can be found in the literature [67–69], however, the potential operative morbidity that is to be expected must also be considered. In the worst case a previously complaint-free but infertile patient – in spite of unsuccessful IVF – may merely experience a marked deterioration in her quality of life. Such scenarios must also be discussed in detail with just those patients who are free of symptoms. More recent data also even additionally question this improvement in fertility, so that the guideline recommendations will probably have to be reassessed on this point [70].

Drug therapy is neither necessary as preparation nor as an adjuvant therapy after the surgical management of DIE. When, in spite of symptomatic DIE – initial or permanent – the decision against surgery is made, the classical hormone therapy options (monophasic oral contraceptives, gestagen monotherapy, levonorgestrel IUP) can then be applied. An effect can only be expected during the therapy so that a life-long continuation is a necessity [1].

A malignant degeneration is rare but possible. DIE can typically develop into endometrioid or clear-cell adenocarcinomas in the pelvis without any clearly associated organs [71]. However, this risk is so low that an indication for treatment or surgery cannot be derived from this alone.

**Endometriosis and Sterility**

**Pathophysiology of involuntary childlessness in cases of endometriosis**

There is a high coincidence between sterility and endometriosis [72], the reasons for this have not yet been clarified. It is estimated that 30–50% of the women with endometriosis are infertile [73]. Mechanical factors due to adhesions of the adnexa with disorders of tube motility or, respectively, the ovum pick-up mechanism have been accepted as reasons for sterility. Further causes may involve disordered immunological dysregulation and also a change of the intra-abdominal milieu, due to cytokines, prostaglandins and macrophages. In an egg cell donation programme patients with endometriosis achieved similar pregnancy rates as those women without endometriosis, however the pregnancy rates in case of transfer of embryos form women with endometriosis was markedly lower. This suggests a decrease in embryo quality in cases of endometriosis [74].

![Fig. 6a and b](image)

**Fig. 6a and b** a DIE of the vesicocervical septum and bladder. b Cystoscopic visualisation with intact urothelium over the finding.

![Fig. 7](image)

**Fig. 7** Situation after local full wall excision of DIE of the bladder.
Drug therapy
A hormone therapy for endometriosis can be performed with pure gestagens (e.g., dienogest, medroxyprogesterone acetate, levonorgestrel, chloromadinone acetate, cyproterone acetate and nomegestrol acetate), with monophasic oestrogen-gestagen combination formulations (especially taken in long cycles), as well as by means of GnRH analogues. In cases of low-degree endometriosis (AFS I and II), however, a metaanalysis of 16 randomised and controlled studies did not reveal an improvement in fertility after drug treatment (GnRH analogues, gestagens) in comparison to placebo or expectative procedure [75]. A postoperative drug therapy with GnRH agonists could not improve the spontaneous pregnancy rates of sterility patients and is thus not recommended [55].

Operative therapy
Just for patients desiring to have children and the suspicion of an endometrial disease surgical diagnosis and histological confirmation with simultaneous hysterectomy and chromopertubation of the Fallopian tubes represent the method of choice. In the course of this treatment if at all possible all endometriosis lesions should be surgically excised or thermally destroyed. Endometriosis cysts should be ablated with cyst bag in an organ-sparing manner. The rate of spontaneous pregnancies is in such cases also higher than for the excision endometriosis lesions in stages AFS I and II [52, 53, 76].

Assisted reproduction
The measures of assisted reproduction in almost all stages of endometriosis seem to be more advantageous as compared with a non-surgical process. In cases of minimal endometriosis the probability of a pregnancy after a wait-and-see procedure for 6 months amounts to 28% [77].

Several prospective controlled studies have shown a higher success rate under ovarian stimulation in combination with intrauterine insemination [77]. For younger patients pregnancies occurred within the first cycle after operative management of endometriosis so that a maximum of 3 insemination treatments should be performed [78].

For endometriosis patients the success rate in the course of an in-vitro fertilisation is up to 50% lower than for tubular sterility [79], furthermore, higher drug doses are needed for the markedly lowered responsiveness to gonadotropins [80]. On the basis of a large metaanalysis it has been shown that the implantation rate for endometriosis patients was markedly lower than that in the control group (12.72 vs. 18.08%) [79]. In case of the recurrence of an extensive endometriosis an assisted reproduction by means of in-vitro fertilisation is superior to renewed operative therapy with regard to the pregnancy rate [81]. In the case of advanced endometriosis AFS III and IV, a stimulation for IVF/ICSI in ultra-long protocol after surgical therapy for endometriosis leads to significantly higher pregnancy rates [82, 83].

In patients with endometriosis and the desire to have children surgical clarification and rehabilitation of endometriosis is considered to be the gold standard. Depending on the degree of severity of the endometriosis, a cycle optimisation in combination with ovarian stimulation and accompanying intrauterine insemination should be preferred although not more than 3 treatment cycles may be carried out.

Patients with severe endometriosis should be treated as early as possible by means of IVF/ICSI, if necessary after stimulation in ultra-long protocol.

Complementary Therapy, Rehabilitation and Follow-Up

For chronic courses of endometriosis with the corresponding symptoms many women experience an alleviation of their complaints and an improvement in their quality of life through the use of complementary therapies such as, e.g., acupuncture, Chinese medicine, homeopathy, phytotherapy, osteopathy, physiotherapy etc. However, at present there are no randomised and controlled studies that confirm an evidence-based effect of these therapies [1].

The need for rehabilitation is often present after extensive surgical interventions or in patients with chronic pain. The aim of rehabilitation should be the restoration of physical, mental and social well-being and should thus be generously offered to this patient collective. An important aspect is, however, also the confrontation with the disease, which often has a chronic course and in part is accompanied by unavoidable restrictions and complaints. Afflicted women should be referred to regional and national self-help organisations and self-help groups. The aftercare should be oriented on the symptoms. The patient’s quality of life is in the forefront of all efforts [1].

In the present article we have topically summarised many aspects of endometriosis with reference to the most recent literature. In particular the less common DIE, a form that is associated with a high morbidity, has been discussed in detail. Also sterility as well as the diagnostic and therapeutic consequences are specifically summarised. Besides further important clinical aspects we have in the present article highlighted the most recent research results, in order to close the gap concerning actual research between scientists and clinicians.

Conflict of Interest

None.

References
8 Sampson JA. Peritoneal endometriosis due to menstrual dissemination of endometrial tissues into the peritoneal cavity. Am J Obstet Gynecol 1927; 14: 422–469

suppresses angiogenesis and vessel maturation in endometriotic le-
let-derived growth factor, but not inhibition of VEGF alone, effectively
endothelial growth factor (VEGF), fibroblast growth factor and plate-
giogenesis in the pathophysiology and therapy of endometriosis. Hum
681
mal vascularization in peritoneal endometriosis. Fertil Steril 1993; 59:
21
Gargett CE, Masuda H. Adult stem cells in the endometrium. Mol Hum
2010; 16: 818–834

Figueira PC, Abrao MS, Kriukin G et al. Stem cells in endometrium and
their role in the pathogenesis of endometriosis. Ann Y Ac Sci
2011; 1221: 10–17

Du H, Taylor HS. Stem cells and reproduction. Curr Opin Obstet Gynecol

Laschke MW, Giebels C, Menger MD. Vascollogenesis: a new piece of the
endometriosis puzzle. Hum Reprod Update 2011; 17: 628–636

Grootbuis PC, Nap AW, Winterhager E et al. Vascular development in endo-
angiogenesis 2005; 8: 147–156

Nisolle M, Casanas-Roux F, Anaf V et al. Morphometric study of the strom-
al vascularization in peritoneal endometriosis. Fertil Steril 1993; 59:
681–684

McLaren J. Vascular endothelial growth factor and endometriotic an-

Laschke MW, Menger MD. In vitro and in vivo approaches to study angi-
genesis in endometriophathy and therapy of endometriosis. Hum Reprod
Update 2007; 13: 331–342

Healy DL, Rodgers PA, Hii L et al. Angiogenesis: a new theory for endome-

Laschke MW, Elitzsch A, Vollmar B et al. Combined inhibition of vascular
endothelial growth factor (VEGF), fibroblast growth factor and plate-
let-derived growth factor, but not inhibition of VEGF alone, effectively
suppresses angiogenesis and vessel maturation in endometriotic le-

Feng D, Welker S, Korbel C et al. Protein kinase CK2 is a regulator of an-

Laschke MW, Menger MD. Anti-angiogenic treatment strategies for the

Anaf V, Simon P, El Nakadi I et al. Hyperalgesia, nerve infiltration and
nerve growth factor expression in deep adenomyotic nodules, perito-
eal and ovarian endometriosis. Hum Reprod 2002; 17: 1895–1900

Alvarez P, Chen X, Hendrich J et al. Ectopic uterine tissue as a chronic


Novella-Maestre E, Herrera S, Vila-Vives JM et al. Effect of antiangiogenic
treatment on peritoneal endometriosis-associated nerve fibers. Fertil
Steril 2012; 98: 1209–1217


Haas D, Wurm P, Shamiyeh A et al. Efficacy of the revised Enzian classi-
fication: a retrospective analysis. Does the revised Enzian classification solve the problem of duplicate classification in rASRM and Enzian? Arch Gynecol Obstet 2013; 287: 941–945


Jansen RP, Russell P. Nonpigmented endometriosis: clinical, laparoscop-

Harrison RF, Barry-Kinsella C. Efficacy of medroxyprogesterone treat-
mament in infertile women with endometriosis: a prospective, random-

Renner SP, Rix S, Boos A et al. Preoperative pain and recurrency risk in patients with peritoneal endometriosis. Gynecol Endocrinol 2010; 26:
230–235

Healey M, Ang WC, Cheng C. Surgical treatment of endometriosis: a pros-
spective randomized double-blind trial comparing excision and ablation. Fertil Steril 2010; 94: 2536–2540

Duffy JM, Arambage K, Correa FJ et al. Laparoscopic surgery for endo-
metriosis. Cochrane Database Syst Rev 2014; 4: CD011031

Hornstein MD, Yuzpe AA, Barry KA et al. Prospective randomized dou-
ble-blind trial of 3 versus 6 months of nafarelin therapy for endome-

Kitawaki J, Kusuki I, Yamanaka K et al. Maintenance therapy with dien-
Biol 2011; 157: 212–216

Strowitzki T, Marr J, Gerlinger C et al. Detailed analysis of a randomized,

Harada T, Momoda M, Taketani Y et al. Dienogest is as effective as intra-
asal busulphan aceite for the relief of pain symptoms associated with endometriosis–a randomized, double-blind, multicenter, con-

Rev 2009; 2: CD004753

Hudelst C, Oberwinkler KH, Singer CF et al. Combination of transvaginal
sonography and clinical examination for preoperative diagnosis of pel-
vic endometriosis. Hum Reprod 2009; 24: 1018–1024


Alborzi S, Motahhan M, Parsaneshzad ME et al. A prospective, random-
ized study comparing laparoscopic ovarian cystectomy versus fenes-


2399–2402

Hudelst C, Keckstein J. [The use of transvaginal sonography (TVS) for
preoperative diagnosis of pelvic endometriosis]. Praxis (Bern) 1999
2009; 98: 603–607

Khong SY, Bignardi T, Luscombe G et al. Learning curve in the detection of ovar-
ian and deep endometriosis by using magnetic resonance: comparison of
rASRM and Enzian? Arch Gynecol Obstet 2014; 281: 216–221


Legard DJ, O’Donnell DJ, Strockbine TM et al. Diagnostic accuracy of physical examination, transvaginal sonography, rectal endoscopic sonography, and magnetic resonance imaging to diagnose deep infiltrating endo-

Saban L, Guerriero S, Suls R et al. Learning curve in the detection of ovar-
ian and deep endometriosis by using magnetic resonance: comparison with surgical results. Eur J Radial 2011; 79: 237–244

Chopin N, Vieira M, Borghese B et al. Operative management of deeply


Tummon IS, Asher LJ, Martin JS et al. Randomized controlled trial of superovulation and insemination for infertility associated with minimal or mild endometriosis. Fertil Steril 1997; 68: 8–12


Pagidas K, Falcone T, Hemmings R et al. Comparison of reoperation for moderate (stage III) and severe (stage IV) endometriosis-related infertility with in vitro fertilization-embryo transfer. Fertil Steril 1996; 65: 791–795
