Intestinal Vaginoplasty for Congenital Absence of the Vagina

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Congenital absence of the vagina is rare and occurs as a result of aplasia of the Müllerian ducts (46,XX) or complete androgen insensitivity syndrome (AIS—46,XY). Both syndromes are associated with normal female external genitalia and these patients are raised as females. These children are usually treated during adolescence with chronic dilatation of the shallow vaginal introitus (pressure tube technique) or by skin graft vaginoplasty (Mclndoe procedure). Neither of these procedures is entirely satisfactory, as both may lead to neovaginal stenosis, inadequate length, poor lubrication, or all three. We have recently modified the operation first described by Baldwin in which a loop of sigmoid colon or small bowel is isolated, closed at one end, and brought down on its vascular pedicle as a neovagina and anastomosed to the hymenal ring. We have performed this operation on four adolescents (mean age, 15 years) and two infants (aged 4 days and 14 months) with excellent results (mean follow-up, 7.5 years for the adolescents and 1.8 years for the infants).

The advantages of a bowel segment in contrast to a skin graft are: (1) minimal likelihood of "poor take" or later contraction because a vascularized epithelial-lined tube is used; (2) patency and depth can be maintained without a mold and with minimal dilatation; (3) spontaneous mucus production matches that of the normal vagina and facilitates sexual intercourse; (4) dyspareunia frequently seen with skin grafts is avoided by the ability of the intestinal segment to withstand local trauma; and (5) the use of an intestinal segment offers the option of performing a bowel interposition vaginoplasty during infancy at the time of surgical correction of more complex associated caudal anomalies. The sigmoid colon is the best choice for interposition vaginoplasty because of size, location, and ease of preserving blood supply.

INDEX WORDS: Vaginal agenesis; Müllerian aplasia; androgen insensitivity syndrome; intestinal vaginoplasty.

CONGENITAL absence of the vagina is rare, with an incidence of 1 in 4,000 to 5,000 women. It occurs as the result of aplasia of the Müllerian ducts (46,XX) or complete androgen insensitivity syndrome (AIS—46,XY). Both syndromes are associated with normal female external genitalia, and the patients are raised as females. Absence of the vagina is usually discovered during adolescence because of amenorrhea or failure to achieve intercourse. The most common form of treatment consists of chronic pressure dilatation of the shallow vaginal introitus by use of a graduated series of rods, the Frank method,1 or by creating a skin graft vaginoplasty around a Styrofoam mold, the McIndoe procedure.2 Neither of these procedures is entirely satisfactory, requiring what most patients regard as unpleasant instrumentation of the vagina with application of dilators or molds for several months. Both procedures may lead to neovaginal stenosis, inadequate length, poor lubrication, or all three. Even temporary noncompliance with the program of dilatation and night-time stent insertion can lead to contraction, scarring, and failure.3

We have recently modified the operation first described by Baldwin4 wherein a loop of sigmoid colon or small bowel is isolated, closed at one end, and brought down on its vascular pedicle as a new vagina and anastomosed to the introitus at the level of the hymenal ring. We have performed this operation on four adolescents (mean age, 15 years) and two infants (aged 4 days and 14 months) with excellent results.

MATERIALS AND METHODS

A summary of the diagnosis, physical findings, and operative treatment is presented in Table 1. The goal of the operation is to fashion a flexible and functional neovagina of bowel readily available from the gastrointestinal tract. This is most effectively accomplished using sigmoid colon as in three of our four adolescent patients, or a segment of ileum as used in the fourth. With respect to the two infants, segments of bowel were used that otherwise would have been discarded: a segment of terminal ileum attached to occum forming the back wall of a bladder duplication in case 5, and a cecal duplication in case 6.

The patient is placed in the lithotomy position with her legs in stirrups or popliteal supports so that the abdomen and vaginal introitus can be prepped in one field. Broad-spectrum antibiotics are begun at the time of anesthesia induction, and a Foley catheter is placed in the bladder. The abdomen and pelvis are approached through a Pfannenstiel incision. In cases of Müllerian agenesis, exploration usually shows separated remnants of uterine horns along with portions of tubes. The ovaries are usually normal in size and position. In cases of AIS (testicular feminization and its variants), the gonads are removed to prevent virilization and obviate the risk of testicular cancer. A 10- to 15-cm length of sigmoid colon is isolated, preserving at least one major artery and vein at one end of the segment along with the accompanying vascular arcade (Fig 1). Depending on the local anatomy of the sigmoid loop, one end or the other is selected for the vascular arcade. The isolated loop on its vascular supply 180° to affect a
Table 1. Intestinal Vaginoplasty for Congenital Absence of the Vagina

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Age at Operation</th>
<th>Diagnosis</th>
<th>Associated Anomalies</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>17 yr</td>
<td>Congenital absence of vagina and uterus; 46,XX (Rokitanski-Kuster-Hanser syndrome)</td>
<td>Absent left kidney; right pelvic kidney</td>
<td>Sigmoid colon vaginoplasty</td>
</tr>
<tr>
<td>2</td>
<td>15 yr</td>
<td>Congenital absence of vagina and uterus; 46,XX (Rokitanski-Kuster-Hanser syndrome)</td>
<td>None</td>
<td>Sigmoid colon vaginoplasty</td>
</tr>
<tr>
<td>3</td>
<td>15 yr</td>
<td>Testicular feminization syndrome; 46,XY</td>
<td>None</td>
<td>Sigmoid colon vaginoplasty, bilateral gonadectomy</td>
</tr>
<tr>
<td>4</td>
<td>14 yr</td>
<td>Testicular feminization syndrome; 46,XY</td>
<td>None</td>
<td>Ileovaginoplasty, bilateral gonadectomy</td>
</tr>
<tr>
<td>5</td>
<td>14 mo</td>
<td>Testicular feminization syndrome; 46,XY</td>
<td>Imperforate anus; duplicated bladder</td>
<td>Ileovaginoplasty; cecal augmentation cystoplasty; bilateral gonadectomy</td>
</tr>
<tr>
<td>6</td>
<td>4 d</td>
<td>Testicular feminization syndrome; Exstrophy of bladder; omphalocele; right clubfoot; cecal duplication</td>
<td>Exstrophy of bladder, omphalocele, right clubfoot; cecal duplication</td>
<td>Colon vaginoplasty (using cecal duplication)</td>
</tr>
</tbody>
</table>

A smoother course to the vaginal orifice with less tension. A satisfactory conduit will result regardless of the direction of peristalsis. We have been able to prepare and position the sigmoid loop in each of these cases, without undue tension on the blood supply.

The continuity of the sigmoid colon or ileum is reestablished with a single layer anastomosis of interrupted silk sutures, and the defects in the mesentery are closed (Fig 3). The hymenal region of the vulva is then incised in a circular or cruciate fashion and the vaginal tract bluntly dissected between the bladder and rectum to the level of the peritoneal reflection in the cul-de-sac. This portion of the procedure can also be done from above downward; specifically, a Hegar dilator can be pushed upward in the rudimentary introitus, and the bulging peritoneum in the cul-de-sac cut down on...
Fig 3. Colocolostomy has been completed and the distal end of the colonic segment has been anastomosed to the opened rudimentary vaginal pit.

Fig 4. The completed vaginoplasty has been loosely packed with Vaseline gauze and the peritoneum closed above the transposed bowel.

RESULTS

The results are outlined in Table 2. There were no operative complications in any of the six intestinal vaginoplasties. Experience with the first three patients (cases 1 to 3, Tables 1 and 2) has been particularly gratifying. Each patient was taught, and easily adapted to, the neovagina dilatation with a Hegar dilator ranging in size from 20 to 26. This might more appropriately be termed a calibration rather than a dilatation in that the neointroitus and colovagina remained generous and supple in each patient. Weekly calibration was continued for 6 months to 1 year, until sexual intercourse made calibration no longer necessary. All three patients report satisfactory and satisfying sexual intercourse: the first two are married and the third has a boyfriend. None of the three require lubrication, and only one reported occasional spotting from mucus production. The fifth patient who underwent an ileovaginoplasty was last seen at age 17, 2½ years after her procedure, at which time she had a tight vaginal vault. She had been instructed in daily dilatation with a no. 20 Hegar dilator. She was not sexually active. Compliance with the dilatation program was poor, and she has been since lost to further follow-up.

The first of the two infants was born with an imperforate anus and initially underwent an ileostomy and mucus fistula, with an 8-cm segment of ileum coming off of a foreshortened colon, consisting of cecum attached to the dome of a neurogenic bladder duplication. She underwent a cecal augmentation cystoplasty at age 14 months, at which time the residual ileum, which had been part of the mucus fistula, was brought down to form a neovagina. She is now 5 years old and undergoes periodic dilatation of the neovagina with a no. 6 Hegar dilator in the office. The vaginal orifice is tight and she will probably require enlargement of the introitus when she reaches adolescence.

The second infant (case 6) was born with multiple congenital anomalies, including exstrophy of the bladder, omphalocele, and a cecal duplication. The latter, at the time of closure of the omphalocele and exstrophy, could be easily mobilized and brought down to fashion a neo-vagina. This seemed to be a
very appropriate use for the segment of duplicated bowel rather than to resect and discard it.

She died at age 16 months of complications of home parenteral nutrition.

DISCUSSION

Congenital absence of the vagina is most commonly caused by Müllerian aplasia, and is frequently called by the eponym Rokitansky-Küster-Hauser syndrome. Müllerian aplasia is usually accompanied by a rudimentary uterus in the form of bilateral and noncannulated muscular buds, normal tubes and ovaries, normal female secondary sex characteristics, and otherwise normal endocrine and cytogenetic evaluations. The differential diagnosis of absence of the vagina in an otherwise normal appearing female includes partial absence of the vagina (usually the lower third) and testicular feminization, or the AIS. The latter may be diagnosed during infancy due to the high incidence of bilateral inguinal hernias. The vagina in these patients will be abnormally shallow or absent all together.

Because of the high incidence (up to 50%) of isolated urinary tract anomalies in patients with congenital absence of the vagina, such as absence of one kidney, horseshoe kidney, pelvic kidney, or duplicated collecting systems, intravenous pyelography or a computed axial tomography scan with intravenous contrast should be done preoperatively. Skeletal malformations are common (up to 12%) and spinal films may show vertebral anomalies. However, the most important aspect of the preoperative evaluation is the establishment of a close doctor-patient relationship such that the surgeon is assured that the patient will cooperate with the use of vaginal dilators and return regularly for follow-up evaluation.

The Frank technique of nonoperative forceful dilatation of the vaginal introitus is most successful in highly motivated individuals with a vaginal dimple of 3 or 4 cm. The McIndoe technique and its variations has been widely applied. Good results can be obtained provided that the graft is complete. Its successful function and result is dependent on a mold being worn for many months in order to overcome the contractile phase of the healing skin graft.

Lubrication is required for satisfactory sexual intercourse and dysparenia has been a frequent complaint. There are now seven reported cases of squamous cell carcinoma arising in split-thickness skin graft vaginoplasties, and two reported cases of adenocarcinoma in segmental bowel vaginoplasties.

Our experience and that of others has been that the sigmoid colon serves as an extraordinarily effective substitute for the vaginal canal. Obtaining adequate length is no problem, and there is no tendency toward contraction, narrowing, or stenosis provided that the bowel segment has an adequate blood supply, and that the anastomosis to the hymenal region is generous. Wearing a stent is unnecessary and dilations, if needed at all, are temporary, infrequent (we recommend once weekly), and well tolerated.

The thick wall of the colon tolerates trauma with less reaction and bleeding than does small bowel or split-thickness skin graft. Bleeding may be intermittent or bothersome during the first month or two, but thereafter rarely occurs. Initially exuberant mucus production by the sigmoid neovagina gradually tapers off over 1 to 2 months and generally has not been a problem. The mucosa atrophies to a mild degree in its new position, and once-a-week douches are helpful to prevent collections of inspissated mucus. Moreover, mucus production is sufficient to keep the neovagina moist, and lubrication before intercourse is unnecessary.

The advantages of a bowel segment vaginoplasty

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**Table 2. Results and Follow-Up of Intestinal Vaginoplasty for Congenital Absence of the Vagina**

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Operation</th>
<th>Duration of Follow-Up</th>
<th>Anatomic</th>
<th>Physiological</th>
<th>Pertinent Postoperative History</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Sigmoid colon vaginoplasty</td>
<td>6 yr</td>
<td>Excellent</td>
<td>Married, normal sexual intercourse</td>
<td>Weekly dilatation for 6 mo, then regular intercourse; needs no lubrication</td>
</tr>
<tr>
<td>2</td>
<td>Sigmoid colon vaginoplasty</td>
<td>12 yr</td>
<td>Excellent</td>
<td>Married, normal sexual intercourse</td>
<td>Weekly dilatation for 1st year, then sporadic; needs no lubrication; occasional spotting</td>
</tr>
<tr>
<td>3</td>
<td>Sigmoid colon vaginoplasty</td>
<td>11 yr</td>
<td>Excellent</td>
<td>Excellent, normal sexual intercourse</td>
<td>Weekly dilatation for 1st year; needs no lubrication</td>
</tr>
<tr>
<td>4</td>
<td>Ileovaginoplasty</td>
<td>2.5 yr</td>
<td>Good</td>
<td>Unknown, unmarried, no sexual intercourse</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Ileovaginoplasty; cecal augmentation cystoplasty</td>
<td>4 yr</td>
<td>Good, narrow introitus</td>
<td>NA</td>
<td>Requires periodic dilatation</td>
</tr>
<tr>
<td>6</td>
<td>Colon vaginoplasty (using cecal duplication)</td>
<td>5 mo</td>
<td>Excellent</td>
<td>NA</td>
<td>Died at home, age 16 mo, suspected pulmonary embolus</td>
</tr>
</tbody>
</table>

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over prolonged forceful dilation of a shallow rudimentary vaginal pit, or split-thickness skin graft inserted over a mold are: (1) minimal likelihood of “poor take” or later contraction because a vascularized epithelial-line tube is used; (2) initial length is no problem and patency and depth can be maintained without a mold and with minimal dilatation; (3) spontaneous mucus production matches that of the normal vagina and facilitates comfortable sexual intercourse; (4) dyspareunia, frequently experienced with skin grafts, is avoided by the greater resistance of the sigmoid segment to local trauma; (5) the operation is straightforward, safe, and unaccompanied by the scarring associated with split thickness skin grafts; and (6) the use of an intestinal segment offers the options of performing bowel interposition vaginoplasty during infancy at the time of surgical correction of more complex associated caudal anomalies. The sigmoid colon is the best choice for interposition vaginoplasty because of size, location, and ease of preserving blood supply. Minimal long-term care is required, although surveillance must be maintained at regular intervals (every 6 months) to watch for adenocarcinoma in the transplanted bowel segment.

REFERENCES