Sacrosinous hysteropexy versus vaginal hysterectomy with suspension of the uterosacral ligaments in women with uterine prolapse stage 2 or higher: multicentre randomised non-inferiority trial

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ABSTRACT

OBJECTIVE
To investigate whether uterus preserving vaginal sacrosinous hysteropexy is non-inferior to vaginal hysterectomy with suspension of the uterosacral ligaments in the surgical treatment of uterine prolapse.

DESIGN
Multicentre randomised controlled non-blinded non-inferiority trial.

SETTING
4 non-university teaching hospitals, the Netherlands.

PARTICIPANTS
208 healthy women with uterine prolapse stage 2 or higher requiring surgery and no history of pelvic floor surgery.

INTERVENTIONS
Treatment with sacrosinous hysteropexy or vaginal hysterectomy with suspension of the uterosacral ligaments. The predefined non-inferiority margin was an increase in surgical failure rate of 7%.

MAIN OUTCOME MEASURES
Primary outcome was recurrent prolapse stage 2 or higher of the uterus or vaginal vault (apical compartment) evaluated by the pelvic organ prolapse quantification system in combination with bothersome bulge symptoms or repeat surgery (n=0). Five serious adverse events were reported during hospital stay. None was considered to be related to the type of surgery.

RESULTS
Sacrosinous hysteropexy was non-inferior for anatomical recurrence of the apical compartment with bothersome bulge symptoms or repeat surgery (n=0, 0%) compared with vaginal hysterectomy with suspension of the uterosacral ligaments (n=4, 4.0%, difference −3.9%, 95% confidence interval for difference −8.6% to 0.7%). At 12 months, overall anatomical recurrences, functional outcome, quality of life, complications, hospital stay, measures on postoperative recovery, and sexual functioning did not differ between the two groups.

CONCLUSIONS
Uterus preservation by sacrosinous hysteropexy was non-inferior to vaginal hysterectomy with suspension of the uterosacral ligaments for surgical failure of the apical compartment at 12 months' follow-up.

TRIAL REGISTRATION
Trialregister.nl NTR1866.

Introduction
Pelvic organ prolapse is a common health problem. The prevalence of such prolapse is as much as 40% in women aged more than 45 years, with millions of women affected worldwide, and the incidence is still rising as a result of aging populations and increasing obesity rates. Pelvic organ prolapse has a negative influence on women’s quality of life and is associated with physical, psychological, and sexual problems.

The lifetime risk for prolapse surgery is 11-20%, and worldwide vaginal hysterectomy is the most commonly performed surgical procedure for uterine prolapse. Perforating a hysterectomy for uterine prolapse is not an evidence based practice and whether or not the uterus should be removed is debatable. Uterus preserving procedures such as vaginal sacrosinous hysteropexy, in which the uterus is attached to the sacrospinous ligament, are becoming more popular. In a recent study we found a trend towards more uterus preservation in the Netherlands, which is in line with more women opting to retain their uterus in case of an equal outcome with hysterectomy. Uterus preservation is thought to be less invasive, and in prospective non-randomised and retrospective cohort studies sacrosinous hysteropexy was as effective as vaginal hysterectomy.
hysterectomy, with a similar rate of recurrence and repeat surgery but with a shorter operating time, less blood loss, faster recovery, and fewer complications. A hysterectomy has known benefits as well: it prevents the development of uterine cancer and stops menstrual bleeding in premenopausal women. After vaginal hysterectomy, however, women may be at increased risk of recurrent prolapse since hysterectomy disrupts the supportive structures of the pelvic floor. To prevent future prolapse of the vaginal vault after hysterectomy additional vault suspension is recommended. Randomised controlled trials comparing uterosacral versus sacrospinous hysteropexy concluded that more research on this subject is needed. We tested the hypothesis that sacrospinous hysteropexy is non-inferior to vaginal hysterectomy with suspension of the uterosacral ligaments for surgical failure after 12 months’ follow-up.

Methods
Study design
A detailed version of the trial protocol has been published previously. In short, all women with uterine prolapse at stage 2 or higher (uterine prolapse 1 cm above the hymen or beyond) requiring surgery were invited to participate. Participants were randomly assigned to sacrospinous hysteropexy or vaginal hysterectomy with suspension of the uterosacral ligaments in a non-blinded multicentre randomised controlled non-inferiority trial. Women with coexisting prolapse of the anterior or posterior vaginal wall, or both were able to participate, and cervical elongation together with uterine prolapse was no reason for exclusion. Concomitant repair of anterior or posterior vaginal prolapse (colporrhaphy) was allowed, including anti-incontinence surgery. We excluded women with previous pelvic floor or prolapse surgery, known malignancy or an abnormal cervical smear test result, a wish to preserve fertility, language barriers, immunological or haematological disorders interfering with recovery after surgery, abnormal ultrasound findings of the uterus or ovaries, or abnormal uterine bleeding, and those who were unwilling to return for follow-up.

All four participating centres were Dutch large non-university teaching hospitals. Centres had to offer both treatment modalities, and we invited them to participate if they were known to perform the interventions in the same standardised manner. All gynaecologists were experienced and had performed a minimum of 20 procedures of each intervention before the start of the trial. As the participating hospitals were teaching hospitals, residents were allowed to perform procedures under direct supervision of the gynaecologist. Gynaecologists and residents of the participating centres assessed the eligibility of patients. The decision to treat uterine prolapse surgically was a shared decision by the woman and her gynaecologist.

We gave the women written information and obtained informed consent before randomisation. The women were randomly allocated in a 1:1 ratio using a web based application with computer generated randomisation tables in blocks of four, stratified by hospital and stage of uterine prolapse. The trial was non-blinded as it was impossible to blind surgeons and women to the allocated surgical procedure. An independent doctor who was not involved in treatment carried out the 12 month follow-up.

Outcome measures
Initially the primary outcome was surgical failure at the 12 month follow-up. As outcome definitions to evaluate prolapse surgery were improved after the start of this trial, we changed the primary outcome during enrolment and before data analysis into the composite outcome measure of recurrent pelvic organ prolapse stage 2 or higher in the apical compartment (uterus or vaginal vault) with bothersome bulge symptoms or repeat surgery for recurrent apical prolapse at the 12 month follow-up. Isala hospital’s medical ethical committee approved this change, and during enrolment we published the protocol in an open access online journal.

We evaluated the original primary outcome, overall anatomical failure (pelvic organ prolapse stage 2 or higher in any compartment), as a secondary outcome together with two additional definitions of surgical failure or success. Firstly, a composite outcome of success, defined as no prolapse beyond the hymen, no bothersome bulge symptoms, and no repeat surgery or pessary use for recurrent prolapse within 12 months. Secondly, overall surgical failure: pelvic organ prolapse stage 2 or higher, pessary use, or repeat surgery for recurrent prolapse in any compartment within 12 months. Other secondary outcomes were functional outcome, quality of life, complications, hospital stay, postoperative recovery, and sexual functioning.

Interventions
We referred to a detailed description of the study interventions to ensure a uniform technique among the surgeons. All women received perioperative antibiotics, prophylaxis against thrombosis, and a bladder catheter according to local hospital protocol. Analgesics were given postoperatively in accordance with each hospital’s protocol. The women were advised to abstain from heavy physical work for six weeks.

Sacrospinous hysteropexy—Vaginal sacrospinous hysteropexy was performed unilaterally to the right sacrospinous ligament. The posterior vaginal wall was
incised and the sacrospinosus ligament accessed through the paraarectal space. Two permanent sutures (Prolene 1.0; Ethicon, Somerville, NJ) were placed under direct vision through the sacrospinosus ligament at least 2 cm from the ischial spine. Additional anterior or posterior vaginal wall repair or incontinence surgery was performed if indicated. The permanent sutures were also placed through the posterior side of the cervix and tightened and the uterus redessed. The posterior vaginal wall was closed with absorbable sutures (Vicryl 2; Ethicon, Somerville, NJ). (Also see www.youtube.com/watch?v=ySSfy2A1_RM and www.youtube.com/watch?v=wjct1r37sTw).

Vaginal hysterectomy—The vaginal wall around the cervix was circumcised. After bladder and bowel dissection the anterior and posterior peritoneum were opened. The uterosacral ligaments—strong supportive ligaments that attach the cervix to the sacrum—were identified, ligated, and transected. The uterus was released in several steps using clamps and sutures. After removal of the uterus, the surgical pedicles were inspected for haemostasis and the adnexa inspected. The peritoneum was closed using a delayed absorbable suture (Vicryl 1.0; Ethicon, Somerville, NJ). Additional vault suspension in this study was performed by suspension of the uterosacral ligaments. Such suspension involves the attachment of the uterosacral ligaments to the vaginal vault, thereby restoring normal support to the apical compartment. Concomitant anterior or posterior vaginal wall repair and anti-incontinence surgery were performed afterwards if indicated.

Measurements and procedures
Gynaecological examination before surgery included pelvic ultrasonography to exclude uterine or ovarian disease, a cervical smear test, and vaginal inspection in 45° semi-upright position for staging pelvic organ prolapse using the pelvic organ prolapse quantification system (POP-Q). This system involves quantitative measurements of various points of the vaginal wall, using the hymen as a reference point. The degree of prolapse of the anterior vaginal wall, posterior vaginal wall, and uterus or vaginal vault is measured in centimetres either above or proximal to the hymen (negative number) or beyond, or distal to the hymen (positive number), with the plane of the hymen being defined as zero. The genital hiatus, perineal body, and total vaginal length are also measured. Based on these measurements, a stage (0-4) is determined for each compartment. The overall stage is equal to the stage of the most severely prolapsed compartment. The women came to the hospital to be examined for pelvic organ prolapse stage at baseline and at six weeks, six months, and 12 months after surgery and annually thereafter until 60 months' follow-up. At the time of the follow-up visits women completed validated health related and disease specific quality of life questionnaires: short form-36, Euroqol 5D, urogenital distress inventory, defecatory distress inventory, and incontinence impact questionnaire. We defined the presence of bothersome bulge symptoms after surgery as a positive answer to any of the following two questions from the urogenital distress inventory: “Do you experience a sensation of bulging or protrusion from the vagina?” and “Do you have a bulge or something falling out that you can see in the vagina?” in combination with a response “somewhat bothered” to “very much bothered” to the question “how much does this bother you?” To assess sexual functioning, we used the 12 item pelvic organ prolapse/urinary incontinence sexual questionnaire, translated from the validated questionnaire but not validated for Dutch language.

During hospital admission and the first six weeks after surgery, the women kept a diary to evaluate postoperative pain (range 0-10), measured by a validated visual analogue scale; analgesics; and postoperative recovery, measured with the validated recovery index-10.

Data were entered and registered using a web based application facilitated by the Dutch consortium for studies in women’s health and reproductivity (www.studies-obsyn.nl).

Statistical analysis
The sample size for this trial was based on the primary outcome. Recurrence rates in the apical compartment after vaginal hysterectomy vary from 0-12%, so that a failure rate including bothersome symptoms and repeat surgery after sacrospinosus hysteropexy of 10% or more might be regarded as high. As we expected a failure rate of 3% based on outcomes of vaginal hysterectomy in a previous randomised study, we set the non-inferiority margin at 7%. This means that when the upper limit of the 95% confidence interval for the estimated difference in recurrence rate after sacrospinosus hysteropexy versus vaginal hysterectomy exceeds 7%, sacrospinosus hysteropexy is inferior to vaginal hysterectomy. Assuming an absolute recurrence rate of 3% in both groups and a two sided α risk of 0.05, with two groups of 94 women each the trial had 80% power with a prespecified non-inferiority margin of 7% to assess non-inferiority of sacrospinosus hysteropexy. Considering a 10% loss to follow-up, we required a 208 women—104 in each study arm.

We assessed study outcomes by intention to treat analysis. In case of missing data on anatomical outcome at 12 months we applied two strategies. For the first by intention to treat analysis, we used the last observation carried forward with data at the six month follow-up visit if available. If these data were not available, we left the women out of the intention to treat analysis. In case of missing data on outcome at 12 months we applied two strategies. For the second by intention to treat analysis we applied conservative imputation for all women with missing data at 12 months (worst case scenario, failure). In the case of missing questionnaires, we obtained information on the presence or absence of bothersome bulge symptoms from the 12 month follow-up visit.

We also carried out a per protocol analysis on the primary and secondary outcomes for anatomical and
Flow of women through study. *Intention to treat: two patients allocated to vaginal hysterectomy (VH) received sacrospinous hysteropexy (SH) and were analysed in the VH group. Data at six and 12 month follow up were missing in one patient after SH and six patients after VH; one patient after VH had recurrent apical prolapse but pelvic organ prolapse quantification (POP-Q) score was missing, this patient was included in the intention to treat-last observation carried forward analysis. †Missed data imputed as failure. ‡Per protocol analysis: two patients did not receive intended treatment. Excluded per protocol analysis: lost for follow-up at 12 months (n=8), missing or incomplete POP-Q score (n=5), and major protocol deviations (n=9); two patients met two criteria to be excluded from per protocol analysis

surgical failure. This analysis included women who completed the entire treatment protocol as originally planned, with availability of the pelvic organ prolapse quantification scores at 12 months and absence of major deviations from the protocol. The Agresti-Coull method was used to calculate 95% confidence intervals for differences in proportions.25 For exploratory purposes we used Fisher’s exact tests and Mann-Whitney U tests to compare proportions and continuous variables between the groups. We used paired sample t tests to compare mean continuous data within groups. After Bonferroni multiple testing adjustment, we considered a P value of less than 0.002 to be significant.32 All statistical analyses were performed with SPSS for windows (version 22.0.0.1).

Patient involvement
No patients were involved in the design and implementation of the study, the dissemination of the results, setting the research question or the outcome measures, or recruitment.

Results
Between 27 November 2009 and 12 March 2012, 208 women were randomly assigned to sacrospinous hysteropexy (n=103) or to vaginal hysterectomy (n=105). The figure shows the flow of women through the study. Baseline characteristics were similar between the groups (table 1) and pelvic measurements and characteristics did not differ at baseline (table 2).

Table 3 presents the results on the primary outcome and the additional definitions of surgical failure. Sacrospinous hysteropexy was non-inferior to vaginal hysterectomy for anatomical recurrence of the apical compartment with bothersome bulge symptoms or repeat surgery for recurrent apical prolapse: sacrospinous hysteropexy 0% (n=0) versus vaginal hysterectomy 4.0% (n=4), difference –3.9% (95% confidence interval –8.6% to 0.7%) for the intention to treat-last observation carried forward approach. Non-inferiority of sacrospinous hysteropexy was also shown in the intention to treat analysis with conservative imputation and the per protocol analysis. The original primary outcome variable of overall anatomical failure occurred in 50% of the women after sacrospinous hysteropexy compared with 44% after vaginal hysterectomy (95% confidence interval for difference –7.4% to 20.1%). No notable differences were found for anatomical recurrences in the different compartments, except for the posterior vaginal wall: sacrospinous hysteropexy 4.4% versus vaginal hysterectomy 16% (95% confidence interval for difference –18.2% to –1.8%).

Table 4 shows the intraoperative and postoperative details of the women, including the secondary outcomes of complication rate and length of hospital stay. Five serious adverse events were reported during hospital stay: two after vaginal hysterectomy and three after sacrospinous hysteropexy. One woman developed paralytic ileus after vaginal hysterectomy. She had also experienced this problem after orthopaedic surgery. She aspirated gastric contents eight days after surgery, developed aspiration pneumonia, and died because of multi-organ failure. The other serious adverse events were atrial fibrillation, which required cardioversion (vaginal hysterectomy); stroke two days after surgery, but with full recovery and no loss of function (sacrospinous hysteropexy); postoperative pneumonia (sacrospinous hysteropexy); and anaphylactic reaction to prophylactic antibiotics before the surgical procedure (sacrospinous hysteropexy); in this last woman the surgical procedure was postponed for several months, without any problems. None of the serious adverse events were judged to be related to the type of surgery.

Tables 5 and 6 provide details on the other secondary outcomes. Functional outcome and quality of life did not differ significantly between the groups (table 5). Postoperative recovery was similar after both interventions, with comparable recovery index-10 scores at 1, 2, 4, and 6 weeks after surgery (table 6). Among the patients who completed the pelvic organ prolapse/urinary incontinence sexual questionnaire before and after surgery, there was significant improvement in scores in both surgical groups (P<0.002 each) but no significant difference in total scores between both interventions (table 6).
Pain scores on the visual analogue scale did not differ notably between both interventions, except for day 14 in favour of hysterectomy. In eight out of nine women who experienced buttock pain, a typically reported problem after sacrospinous hysteropexy, the pain resolved (visual analogue scale score ≤2) spontaneously within the first six weeks. One woman underwent suture cutting and vaginal hysterectomy after four months because of persistent pain localised at the place of the sacrospinous hysteropexy sutures. After this procedure she was free of symptoms.

Fourteen protocol deviations occurred: two women received sacrospinous hysteropexy instead of vaginal hysterectomy owing to technical difficulties during surgery (crossovers). In one woman allocated to vaginal hysterectomy, laparoscopic cystectomy before vaginal hysterectomy showed intra-abdominal adhesions, and an abdominal hysterectomy was performed. In two women an exclusion criterion was ignored before randomisation: one woman had had previous pelvic floor surgery (repair of posterior vaginal wall prolapse) and another had primary sclerosing cholangitis with thrombocytopenia. Three women who were assigned to sacrospinous hysteropexy had abnormal smear test results and were treated with electro-surgical excision (n=2) during prolapse surgery or follow-up. In six of 102 women (6%) apical suspension after vaginal hysterectomy was performed using a McCall procedure instead of suspension of the uterosacral ligaments. The other 96 women underwent suspension of the uterosacral ligaments (94%). According to the intention to treat principle, we included these women in the intention to treat analysis, with all women analysed as randomised. In the per protocol analysis we excluded these women, except for those classified as crossovers (n=2), as the primary outcome was related to efficacy and these women had no other protocol deviations, and those with abnormal smear test results (n=3), as this was regarded as a minor protocol deviation.

Residents performed three of the procedures in the sacrospinous hysteropexy group (3%) and 19 in the vaginal hysterectomy group (18%). Overall surgical failure rate (recurrent prolapse, pessary use, or repeat surgery) did not differ significantly by gynaecologist or resident (91 failures out of 180 procedures (50.6%) versus nine failures out of 19 procedures (47.4%), respectively, P=0.81). Endometrial cancer was diagnosed during follow-up in one woman in the sacrospinous hysteropexy group (1.0%), and a laparoscopic hysterectomy was performed.

Urinary retention, defined as more than 150 mL urine retention after removal of the catheter, was similar between the groups (table 4). Affected women received a transurethral catheter or were instructed to perform clean intermittent self catheterisation. In all women spontaneous micturition was achieved after a maximum length of catheterisation of 40 days (median 5.0 days, range 1-40 days). Subsequent surgical treatment for stress urinary incontinence after 12 months was carried out in 1 of 102 women (1.0%) after sacrospinous hysteropexy and 4 of 102 women (3.9%) after vaginal hysterectomy (P=0.37).

**Discussion**

Treatment with sacrospinous hysteropexy was non-inferior to vaginal hysterectomy with suspension of the uterosacral ligaments for surgical failure of the apical compartment in both an intention to treat analysis and a per protocol analysis. We found no notable differences
suggest that the highest risk of prolapse surgery after hysterectomy is in the first two postoperative years. Furthermore, not only is long term surgical outcome important but also short term secondary outcomes, such as complication rate and postoperative recovery. The women consented to follow-up for 60 months after surgery, and these data will be further analysed.

After vaginal hysterectomy the ligatures of the uterosacral ligaments were to be sutured to the vaginal vault to aid in long term vaginal support. However, the protocol was ignored in six women and the McCall procedure was performed instead of suspension of the uterosacral ligaments. Both procedures rely on the uterosacral ligaments for support of the vaginal apex but are considered different treatment modalities for suspension of the vaginal vault, which could have led to differences between treatments. As far as we know, strong evidence on the best technique for vault suspension after vaginal hysterectomy is lacking in the literature, and a recent published trial found similar outcomes after suspension of the uterosacral ligaments and sacrospinous fixation for apical prolapse. In the per protocol analysis these women were excluded, but this did not alter the conclusions.

Another limitation might be that residents were allowed to perform sacrospinous hysteropexy or vaginal hysterectomy under direct supervision of a gynaecologist because of their training position. Surgery by residents may have led to variation in procedures. In the hysterectomy group more procedures were performed by residents. No statistically significant difference was found in surgical failure rate in women who underwent surgery by either gynaecologist or resident, but the higher number of procedures performed by residents could have contributed to a longer operation time in the vaginal hysterectomy group. Surgery performed by residents may improve the generalisability of the trial findings as their involvement in treatment of pelvic organ prolapse is common in Dutch urogynaecological practice.

Comparison with other studies
The anatomical outcome after sacrospinous hysteropexy or vaginal hysterectomy under direct supervision of a gynaecologist because of their training position. Surgery by residents may have led to variation in procedures. In the hysterectomy group more procedures were performed by residents. No statistically significant difference was found in surgical failure rate in women who underwent surgery by either gynaecologist or resident, but the higher number of procedures performed by residents could have contributed to a longer operation time in the vaginal hysterectomy group. Surgery performed by residents may improve the generalisability of the trial findings as their involvement in treatment of pelvic organ prolapse is common in Dutch urogynaecological practice.

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in combination with bothersome bulge symptoms or repeat surgery for recurrent apical prolapse. In our opinion, this composite outcome measure is more clinically relevant than outcome in terms of objective pelvic organ prolapse quantification scores alone. Barber and colleagues reported on different definitions of success after surgery for pelvic organ prolapse in 2009. Treatment success varied widely depending on the definition used, but definitions that included the absence of vaginal bulge symptoms had the strongest relation with the women's assessment of overall improvement and treatment success. Furthermore, the authors concluded that the hymen is an important cut-off point for development of symptoms. As new trials probably will use these updated outcome definitions, we also analysed our data using the hymen as an anatomical threshold and also used the composite outcome measure of Barber and colleagues, making this trial in the future comparable to that of others.

Although the presence of posterior vaginal wall prolapse in both groups was similar before surgery, more repairs of the posterior vaginal wall (colporrhaphies) were performed in the vaginal hysterectomy group. The surgeons were free to decide on concomitant surgery and in general this was decided during surgery. One explanation might be that the surgeons thought that the more dorsal axis of the vagina after sacrospinous hysteropexy already protected against a recurrent prolapse of the posterior vaginal wall. Despite the hysterectomy already protected against a recurrent prolapse of the posterior vaginal wall, more anatomical recurrences of the posterior compartment occurred after hysterectomy with suspension of the uterosacral ligaments, which supports this view. The risk for recurrent prolapse of the anterior vaginal wall after sacrospinous hysteropexy is often discussed. We found no difference in occurrence of prolapse of the anterior vaginal wall. This is in line with previous studies: a retrospective study by Smilen and colleagues found that the occurrence of prolapse of the anterior vaginal wall was not altered by sacrospinous hysteropexy and the randomised study performed by Dietz and colleagues did not find a higher rate of prolapse of the anterior vaginal wall after sacrospinous hysteropexy (51%) compared with vaginal hysterectomy (64%) after one year.

Reoperation rates for (recurrent) pelvic organ prolapse did not differ. Two women ended up undergoing hysterectomy after uterine preservation. In one woman this was because of persistent buttock pain. The overall rate of buttock pain after sacrospinous hysteropexy in our study (9%) is in line with that of other studies and in most women the pain resolved spontaneously. Pre-operative counselling should include information about the potential risk of buttock pain. Endometrial carcinoma was found in one woman during follow-up (1%). A previous retrospective analysis of disease findings after prolapse surgery with hysterectomy showed premalignant or malignant abnormalities in 17 of 644 patients (2.6%). In the study, two women (0.3%) had endometrial cancer diagnosed. Because of the low incidence and the early diagnosis of endometrial cancer owing to blood loss, we believe that future risk of malignancy should not be regarded as a valid reason for removal of the uterus.

Clinical implications and future research
Uterus preservation has gained popularity among gynaecologists and patients during the past years. A recent trial among 213 women from multiple study sites throughout the United States found that 36% of the women preferred uterus preservation, 20% preferred hysterectomy, and 44% had no preference, assuming equal outcomes after both procedures. Another preference study among 100 women showed that 60% would decline hysterectomy if an equally efficacious alternative was available. This trial provides evidence for sacrospinous hysteropexy being such an alternative and therefore this study has important implications for clinical practice. Women who want to avoid hysterectomy and preserve their uterus can be reassured that sacrospinous hysteropexy was equally effective as vaginal hysterectomy after short term follow-up. However, longer follow-up is necessary, and also randomised controlled trials comparing other uterus preserving procedures are needed.

Conclusions
Based on analysis of 12 months’ follow-up, we conclude that sacrospinous hysteropexy is non-inferior to vaginal hysterectomy with suspension of the uterosacral ligaments for recurrent prolapse of the apical
compartment with bothersome bulge symptoms or repeat surgery for recurrent apical prolapse. Overall anatomical outcome, functional outcome, hospital stay, complications, postoperative recovery, and sexual functioning did not differ.

We thank the staff at the participating centres for their assistance with study enrolment. We also thank N van Rijn for help with data collection and administrative support.

Contributors: RJD, JdB, MEV, and HWFvE conceived and designed the study. RJD, JdB, JS, and HWFvE acquired the data. RJD, JI, and KBK carried out the statistical analysis. RJD and HWFvE obtained funding

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### Table 5 | Functional outcome and quality of life after sacrospinous hysteropexy and vaginal hysterectomy at 12 month follow-up. Values are medians (interquartile ranges) of domain scores unless stated otherwise

<table>
<thead>
<tr>
<th>Domains</th>
<th>Before surgery</th>
<th>12 months after surgery</th>
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<tbody>
<tr>
<td></td>
<td>Sacrospinous hysteropexy (n=101)</td>
<td>Vaginal hysterectomy (n=104)</td>
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<tr>
<td>Urogenital distress inventory†:</td>
<td></td>
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<tr>
<td>Overactive bladder</td>
<td>0 (0-44)</td>
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<td>Urinary incontinence</td>
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<td>Obstructive micturition</td>
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<td>17 (0-33)</td>
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<tr>
<td>Pain</td>
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<td>Defecatory distress inventory†:</td>
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<td>Obstruction</td>
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<td>Obstructive defecation</td>
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<td>Incontinence</td>
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<td>Incontinence impact questionnaire‡:</td>
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<td>Mobility</td>
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</tr>
<tr>
<td>Social functioning</td>
<td>94 (75-100)</td>
<td>88 (75-100)</td>
</tr>
<tr>
<td>Role limitations physical</td>
<td>75 (6-100)</td>
<td>100 (50-100)</td>
</tr>
<tr>
<td>Role limitations emotional</td>
<td>100 (75-100)</td>
<td>100 (100-100)</td>
</tr>
<tr>
<td>Mental health</td>
<td>84 (72-92)</td>
<td>84 (72-88)</td>
</tr>
<tr>
<td>Vitality</td>
<td>70 (50-80)</td>
<td>70 (55-80)</td>
</tr>
<tr>
<td>Bodily pain</td>
<td>78 (59-100)</td>
<td>80 (67-100)</td>
</tr>
<tr>
<td>General health perception</td>
<td>75 (55-85)</td>
<td>75 (61-85)</td>
</tr>
<tr>
<td>Health change</td>
<td>50 (25-50)</td>
<td>50 (50-50)</td>
</tr>
</tbody>
</table>

All patients were analysed as allocated.

*P value for exploratory purposes: Mann-Whitney U test of sacrospinous hysteropexy versus vaginal hysterectomy.
†0 = no symptoms or not bothersome to 100 = most bothersome symptoms.
‡0 = best quality of life to 100 = worst quality of life.
§0 = worst quality of life to 100 = best quality of life.

### Table 6 | Postoperative recovery and sexual functioning after sacrospinous hysteropexy and vaginal hysterectomy at 12 month follow-up. Values are means (standard deviations) unless stated otherwise

<table>
<thead>
<tr>
<th>Time point of assessment</th>
<th>Sacrospinous hysteropexy</th>
<th>Vaginal hysterectomy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of women</td>
<td>Mean (SD) score</td>
</tr>
<tr>
<td>Recovery index-10†:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Week 1</td>
<td>99</td>
<td>32 (7)</td>
</tr>
<tr>
<td>Week 2</td>
<td>100</td>
<td>34 (7)</td>
</tr>
<tr>
<td>Week 4</td>
<td>98</td>
<td>36 (7)</td>
</tr>
<tr>
<td>Week 6</td>
<td>98</td>
<td>38 (6)</td>
</tr>
<tr>
<td>PISQ-12‡:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total score at baseline</td>
<td>56</td>
<td>33 (6)</td>
</tr>
<tr>
<td>Total score at 12 months</td>
<td>49</td>
<td>37 (5)§</td>
</tr>
</tbody>
</table>

PISQ-12= pelvic organ prolapse/urinary incontinence sexual questionnaire.

*P value for exploratory purposes: independent samples t test of sacrospinous hysteropexy versus vaginal hysterectomy.
†10 item questionnaire measuring postoperative recovery on 5 point Likert scale. Summary scale score ranges from 10 to 50, where 50 indicates perfect recovery.
‡Total scores range from 0, which represents poorest sexual function, to 48 best sexual function.
§Not shown: paired sample test baseline score and follow-up score (sacrospinous hysteropexy P < 0.002 and vaginal hysterectomy P < 0.002).
and supervised the study and are the guarantors. RDJ, JdB, and HWFvE provided administrative, technical, or material support. All authors analysed and interpreted the data, drafted the manuscript, critically revised the manuscript for important intellectual content, had full access to all of the data in the study, and take responsibility for the integrity of the data and the accuracy of the data analysis.

**Funding:** This study received an unrestricted grant from the Isala research foundation. The funding source had no role in the design and conduct of the study; collection, management, analysis, or interpretation of the data; or preparation, review, or approval of the manuscript.

**Competing interests:** All authors have completed the ICME uniform disclosure form at www.icmje.org/coi_disclosure.pdf and declare: JdB: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf and declare: JdB, AF, HW: Financial support from medical device and pharmaceutical companies. MV: has received honoraria and other potentially relevant funding from medical device and pharmaceutical companies. All other authors declare no competing interests.

**Ethical approval:** This study was approved by the medical ethical committee of the Isala hospital (MEC 09-625) and the local ethical committees of the participating centres, in accordance with the Declaration of Helsinki.

**Data sharing**

**Declaration of Helsinki.**

**Committees:** The study was approved by the local ethical committees at the participating centres, in accordance with the Declaration of Helsinki.

**Patient consent:** All patients gave written informed consent for their participation in the study.

**Randomization:** We used random block randomization sequences provided by the Cochrane Collaboration Centre for Health Economics, Evidence and Technology Assessment, with block sizes 100. Randomization was stratified on the basis of surgical teams and random numbers were generated in advance. Balancing of the distribution of potential confounders across surgical teams was achieved by the use of an independent statistician who was not involved in the study.

**Sample size calculation:** We based our sample size on a power calculation for a 25% non-inferiority margin. Assuming that about 5% of patients undergoing vaginal hysterectomy would undergo vaginal vault prolapse repair within 1 year after surgery, we estimated that study of 761 patients would be needed to achieve 80% power for non-inferiority. A non-inferiority margin of 5% was chosen to ensure that the surgery would be safe (i.e., at a maximum of 10% of patients in the vaginal hysterectomy group).

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