Objective: The aim of this study was to analyze quantified displacements of the posterior vaginal wall (PVW) on dynamic magnetic resonance imaging (MRI), which may generate hypotheses for the detailed mechanisms that underlie the development of posterior vaginal prolapse.

Methods: Pelvic dynamic MRI scans were obtained for 12 women with normal vaginal structure (stage 0) and 62 women with 4 consecutive stages (1–4) of posterior vaginal prolapse. Structural locations (apex vagina, distal vagina, and mid–perineal body [PB]) and equidistant points along the PVW (points 4–6 were considered as midvagina) were identified, and PVW length, straight distance of PVW, levator ani parameters (levator hiatus length [LHL], levator hiatus width [LHW], levator plate angle, anorectal angle, and M line [ML]), urogenital hiatus, and prolapse diameter were measured at rest and maximal Valsalva, respectively. The displacement of these measurements was obtained.

Results: From stage 0 to 2, the variables LHL, LHW, levator plate angle, anorectal angle, and ML increased gradually, but midvagina, distal vagina, and mid-PB were the opposite. From stage 2 to 3, apex vagina, midvagina, distal vaginal, mid-PB, LHL, LHW, and ML raised rapidly and peaked at stage 3, then declined at stage 4. In addition, the correlation coefficients between each measurement from stage 2 to 3 were statistically higher than those from stage 0 to 2.

Conclusions: Quantified displacements of the PVW and its supporting structure were shown on dynamic MRI, and the mechanical mechanisms were hypothesized regarding the interaction between pressure and the supporting force contributing to the deformation of the PVW and the supporting structures.

Key Words: dynamic magnetic resonance imaging, hypothesis, mechanical mechanism, posterior vaginal prolapse, support structure

S tatistics show that each year more than 200,000 operations are performed to repair vaginal prolapse in the United States alone.¹ Repair of posterior vaginal prolapse (PVP) was consisted in 87% of all pelvic surgeries,² and literature indicates that 25% to 29% of patients will require a second surgical intervention.³,⁴ However, posterior pelvic floor problems were actually neglected in urogynecology.⁵ Although the reason of impairment of pelvic floor is considered universally, the interaction of the pressure and the support force results in the deformation of posterior vagina wall and the supporting structures as the PVP progression has not been described; thus, the underlying mechanism of PVP remains unclear.

At present, dynamic magnetic resonance imaging (MRI) has established as a multiplanar global evaluation method for assessing pelvic contents, and the treatment of pelvic floor dysfunction is increasingly dependent on preoperative imaging.⁶ Existing MRI has indicated that the relative locations of the perineal structures and the apex vagina are more caudally positioned in the posterior vaginal wall (PVW) than in normal conditions at maximal Valsalva state.⁷ Besides, there are clear differences in movement along the length of the anterior and PVW when compared between a resting state and maximum Valsalva.⁸ However, static images do not provide information regarding the conduction of pressure in the pelvic region, and there are no possible methods to test and verify at present.

In this study, we conducted a detailed analysis of the physical displacement of pelvic floor structures in patients with progressive degrees of PVP as measured via dynamic MRI. Based on observed displacements of portions of the PVW under Valsalva loading, we have made several inferences regarding possible mechanisms that underlie the development of PVP. We believe that observing the correlation between displacement of one structure relative to another may provide insight regarding how the interaction between pressure and supporting force causes the deformation of PVW and the supporting structures as prolapse progresses.

MATERIALS AND METHODS

Sixty-two women with rectocele-type PVP and 12 volunteers with normal vaginal support were selected from the urogynecology clinic of the Fuzhou General Hospital, Fuzhou, Fujian Province, China. All patients with PVP had the PVW extended lower than the most dependent part of the anterior wall or cervix. All subjects received gynecology examinations, completed the Pelvic Organ Prolapse Quantification System (POP-Q), and underwent dynamic MRI. According to POP-Q staging, women with PVP were divided into 4 groups: stages 1, 2, 3, and 4, and women defined as stage 0 were healthy, nulliparous volunteers without PVP. Women who had histories of previous hysterectomy or other surgeries for pelvic floor disorders were excluded. Ethical approval was provided by the hospital's ethics committee, and informed consent was obtained from all participants.

With subjects in the supine position, MRI of the pelvic cavity was performed using a Siemens Magnet Trio 1.0 T System (Siemens, Munich, Germany). Three-dimensional (3D)-T2 sequences were used to obtain resting-state MRI scans in the sagittal, coronal, and axial planes. The parameters used for 3D-T2 sequences were as follows: repetition time = 1600 milliseconds, time to echo = 97 milliseconds, field of view = 400 cm, and slice thickness 1.0 mm, interleaved with no gap. Subjects were instructed to carry out a maximal Valsalva maneuver for approximately 15 seconds in order to acquire images of PVW protrusion using a fast-spin proton density technique. Each subject was scanning at least 3 times repeatedly, and it was ensured that the maximal extent of prolapse seen during clinical examination was
reproduced in the scanner. General structural identifications were analyzed by the same radiologist and urogynecologist, who were blinded to the clinical status of the examined patient.

Using the analogical method in previous reports, lociations with numbers 1 through 9 equidistantly for PVW and mid–perineal body (PB) were designated (locations 4–6 were considered midvagina); the straight-line distance of PVW (between apex vagina and distal vagina), PVW length, urogenital hiatus (UH), and levator ani (LA) diameters (levator hiatus length [LHL], levator plate angle [LPA], anorectal angle [ARA], and M line [ML]) were determined. We also calculated the parametric estimation of the movement value for the midvagina. The variation of movement angle of each location between the direction of movement and the long axis of the body was also measured from the resting state to maximum Valsalva (Fig. 1).

Data are exhibited as means ± SD and were tested for normality using the Kolmogorov-Smirnov test. Continuous variables were analyzed using analysis of variance (ANOVA) combined with a post hoc multiple-comparisons test. Correlation analysis was performed by determining Spearman correlation coefficients. P = 0.05 was regarded as the level of statistical significance. All data were analyzed with SPSS software IBM SPSS Statistics 21 (SPSS Inc, Chicago, Ill).

RESULTS

Demographic data for all subjects are shown in Table 1. Among the study participants, there were no significant differences in age, weight, body mass index, vaginal parity, history of forceps midwifery, and postmenopause. Statistically significant differences were found among the 5 groups in terms of point D and point Bp on the clinical POP-Q physical examination.

The mean measured displacement values of each group for PVW support systems from resting state to maximum Valsalva are shown in Table 2. Landmarks of vaginal support (level I represents apex vagina; level II, midvagina; level III, distal vagina and mid-PB), LHL, LHW, ML, UH, and 1 to 9 locations along the...
PVW exhibited higher displacement in stage 3 than other stages, and the variation of LP A and ARA climbed as the stage increased (Figs. 2 and 3A).

In addition, the entire PVW shifted downward; the movement angle for the upper and lower portions of PVW was posterior toward the rectum and slightly caudal, whereas the middle portions of PVW moved ventral. The vaginal wall displacement direction differed among the 5 groups (Fig. 3B). The result of the movement angle is shown in Table 3, and movement angles of wall locations 5 to 8 were significantly different between stages 3 to 4 and stages 0 to 2.

The correlations of the measurements were also analyzed (Table 4). While the variations of the LA diameters were moderately correlated with UH from stage 0 to 2, the correlation coefficient with midvagina was weak; nevertheless, distal vagina and mid-LA showed no correlation. Then from stage 2 to 3, there were relationships between each other among apex vagina, midvagina, distal vagina, LA diameters, and UH.

The correlations between degree of prolapse and LP A, ARA, length of PVW, and PD were 0.405 (P < 0.001), 0.352 (P < 0.01), 0.419 (P < 0.001), and 0.572 (P < 0.001), respectively, but there was no correlation with straight-line distance of PVW.

**DISCUSSION**

To our knowledge, we first presented a possible mechanism of PVP based on the quantified displacements from the perspective of mechanical equilibrium and described a detailed procedure for how the pressure and supporting force interaction contributes to the formation of the PVW and the supporting structures in the progressive degrees of PVP. To better demonstrate and comprehend the hypothesis, the continuous progress was segmented artificially and described as follows.

Here, a basic structural paradigm was needed to help guide discussion: the uterosacral-cardinal ligament and the uterus are compared with “slack cord” and “boat,” respectively; uterus position is not decided by the ligaments tethering the boat to the dock primarily, but by the surrounding structures such as “water.” Linear relationship between pressure and displacement could also help to illustrate the hypothesis.

From stage 0 to 2, because of the curving model and physiologic elastic range, the pull of uterosacral-cardinal ligament was considered zero, despite the incremental displacement of apex vagina. Thus, the pressure difference directly exerted on LA and the diameters of LA increased. When the LA appeared to block

**TABLE 2. Variation of Various Clinical Parameters Related to Vaginal Prolapse**

| Landmark           | Stage 0 (n = 12) | Stage 1 (n = 18) | Stage 2 (n = 24) | Stage 3 (n = 11) | Stage 4 (n = 9) |
|--------------------|-----------------|-----------------|-----------------|-----------------|----------------|
| Apex vagina, mm    | 25.33 ± 16.98   | 27.21 ± 13.17   | 30.93 ± 16.85   | 42.65 ± 16.80   | 30.33 ± 15.62  |
| Midvagina, mm      | 18.34 ± 10.20   | 16.30 ± 8.48    | 15.61 ± 10.10   | 24.95 ± 13.91   | 19.93 ± 9.83   |
| Distal vagina, mm  | 15.00 ± 8.98    | 13.46 ± 9.07    | 12.16 ± 8.82    | 22.73 ± 14.16   | 11.76 ± 3.38   |
| Mid-PB, mm         | 26.19 ± 7.80    | 23.47 ± 9.47    | 22.17 ± 8.82    | 36.96 ± 10.54   | 30.01 ± 5.60   |
| LHL, mm            | 5.30 ± 3.89     | 7.18 ± 3.11     | 9.43 ± 6.23     | 12.47 ± 7.14    | 7.17 ± 6.68    |
| LHW, mm            | 3.79 ± 2.39     | 3.89 ± 3.26     | 9.51 ± 7.22     | 13.48 ± 9.56    | 6.40 ± 9.73    |
| ML, mm             | 3.98 ± 3.87     | 5.49 ± 5.12     | 6.39 ± 5.18     | 12.66 ± 9.78    | 6.83 ± 6.61    |
| LPA, °             | 9.48 ± 3.93     | 10.68 ± 6.24    | 10.95 ± 7.06    | 16.76 ± 5.72    | 18.20 ± 7.95   |
| ARA, °             | 8.76 ± 4.47     | 9.04 ± 5.44     | 12.52 ± 5.72    | 14.00 ± 8.85    | 16.99 ± 10.12  |
| PVW length, mm     | −8.01 ± 11.20   | 6.97 ± 17.30    | −0.36 ± 22.62   | 14.24 ± 26.49   | 36.41 ± 20.40  |
| Straight distance of PVW, mm | −3.87 ± 8.81 | −7.21 ± 5.81 | −8.27 ± 6.10 | −5.41 ± 7.40 | −3.74 ± 11.76 |
| UH, mm             | 5.13 ± 6.07     | 5.47 ± 3.34     | 5.04 ± 4.65     | 7.92 ± 6.29     | 2.63 ± 3.66    |
| Prolapse diameter, mm | 30.07 ± 9.95 | 36.01 ± 7.87   | 37.20 ± 9.22    | 46.09 ± 10.32   | 57.89 ± 10.53  |

Values are reported as mean ± SD. Midvagina indicates mean displacement of points 4 to 6.

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most of the pressure, the rest applied to levels II and III supports became smaller; thus, the displacement of midvagina, distal vaginal, and mid-PB became less, and the deformation of the lower two-thirds of PVW was not obvious. Besides, the displacement of mid-PB was greater than that of midvagina and distal vagina, which means PB gets more stress compared with two others. Jing et al.\textsuperscript{17} also reported that the muscle near the PB region takes the greatest strain.

From stage 2 to 3, the impairment of pelvic floor continued to increase, the resistance of LA was no longer sufficient, and so the vaginal support structure appeared to initiate a suspending and sustaining function. At this moment, although the “slack”

![FIGURE 3. A, Mean magnitude (in mm) of displacement measured at 9 locations along the PVW. Error bars represent SEs, whereas asterisks (*) indicate statistically significant differences of the locations at which displacement occurs (P < 0.05). B, Angle of displacement (degrees from the vertical) in normal women and women with different degrees of PVP.](image)

| TABLE 3. Movement Angle (°) (Degrees From the Vertical) in Normal Women and Women With Different Stages of PVP |
|---|---|---|---|---|---|---|
| Point | Stage 0 (n = 12) | Stage 1 (n = 18) | Stage 2 (n = 24) | Stage 3 (n = 11) | Stage 4 (n = 9) | ANOVA P |
| 1 | −10.7 ± 14.1 | 0.4 ± 9.1 | 9.7 ± 8.9 | 22.5 ± 9.8 | 30.3 ± 7.9 | 0.124 |
| 2 | −4.5 ± 15.8 | 7.5 ± 9.7 | 11.0 ± 8.6 | 9.4 ± 10.5 | 14.5 ± 9.3 | 0.840 |
| 3 | −18.0 ± 12.7 | 6.3 ± 9.3 | 9.1 ± 9.2 | 11.8 ± 10.3 | 17.6 ± 12.7 | 0.290 |
| 4 | −13.0 ± 13.5 | 21.5 ± 7.5 | −4.9 ± 8.4 | 11.4 ± 11.0 | −15.6 ± 11.7 | 0.058 |
| 5 | −20.6 ± 12.4* | 20.7 ± 6.4\textsuperscript{1} | 0.5 ± 6.6\textsuperscript{2} | 8.0 ± 9.3\textsuperscript{3} | −30.9 ± 10.4\textsuperscript{4} | 0.001 |
| 6 | −18.9 ± 10.6 | 21.4 ± 5.4\textsuperscript{1} | −5.7 ± 6.8\textsuperscript{3} | 20.7 ± 9.3\textsuperscript{4} | −33.8 ± 7.5\textsuperscript{5} | 0.000 |
| 7 | −1.3 ± 13.9 | 16.4 ± 6.1\textsuperscript{1} | −1.6 ± 8.6\textsuperscript{3} | 22.6 ± 9.6\textsuperscript{4} | −15.3 ± 6.8\textsuperscript{5} | 0.047 |
| 8 | −11.8 ± 10.2* | 28.1 ± 5.3\textsuperscript{1} | −1.1 ± 7.1\textsuperscript{5} | 25.2 ± 8.7\textsuperscript{5} | −4.5 ± 7.4\textsuperscript{5} | 0.001 |
| 9 | −12.9 ± 17.4 | 30.0 ± 9.7 | 14.1 ± 10.4 | 27.5 ± 11.2 | 10.2 ± 10.6 | 0.147 |

Values are reported as mean ± SD. P values were determined using ANOVA for comparisons of means and least significance difference test for parity.

Movement angle is the angle between the direction of movement and the long axis of the body. Positive to the dorsal, negative to the ventral.

\*Pairwise: stage 3 versus stage 0 (P < 0.05).
\textsuperscript{1}Pairwise: stage 4 versus stage 1 (P < 0.001).
\textsuperscript{2}Pairwise: stage 4 versus stage 2 (P < 0.01).
\textsuperscript{3}Pairwise: stage 4 versus stage 3 (P < 0.05).
\textsuperscript{4}Pairwise: stage 3 versus stage 2 (P < 0.05).
cord transformed into “tighten”\textsuperscript{15} and started to resist pressure, it was still inadequate for contending against the growing pressure difference. The remaining pressure passed down to levels II and III support; thus, displacement of vaginal supports and LA diameters ascended and peaked at stage 3 with the similar movement circle. The vaginal wall also participated in maintaining resistance by enlarging effective stress area and altering shape, and length of PVW was related to degree of prolapse moderately. When vaginal support resisted powerless, PVW deformed violently, the entire PVW shifted downward, and the movement circle. The vaginal wall also participated in main-
In the current phase, we found that correlation coefficient of level I versus lower elements (levels II and III and UH) was larger than LA versus lower elements at stage 0 to 2 universally. In addition, the displacement vector of apex vagina exceeded others; thus, apical support withstood the largest portion of loads and was the one closest to PVP. Haylen et al\textsuperscript{13} also reported midvagina slack was caused by the descent of the apex vagina, and Rooney et al\textsuperscript{19} found a significant correlation between apex vagina and slack was caused by the descent of the apex vagina, and Rooney et al\textsuperscript{19} found a significant correlation between apex vagina and formation path of PVW were in agreement with a previous report by Lewicky-Gaupp et al.\textsuperscript{7}

TABLE 4. Correlation Coefficients Between Different Levels of Vaginal Support, LA, and UH

| Stage | Apex Vagina | Midvagina | Distal Vagina | UH |
|-------|-------------|------------|---------------|----|
|       | Stage 1 | Stage 2 | Stage 3 | Stage 0 | Stage 1 | Stage 2 | Stage 3 | Stage 0 | Stage 1 | Stage 2 | Stage 3 | Stage 0 | Stage 1 | Stage 2 | Stage 3 | Stage 0 |
| LHL   | 0–2   | r = 0.309 | P = 0.023  | r = 0.125 | P = 0.368 | r = 0.567 | P = 0.000 |
|       | 2–3   | r = 0.639 | P = 0.000  | r = 0.368 | P = 0.030 | r = 0.567 | P = 0.000 |
| LHW   | 0–2   | r = 0.219 | P = 0.111  | r = 0.065 | P = 0.638 | r = 0.317 | P = 0.02 |
|       | 2–3   | r = 0.582 | P = 0.000  | r = 0.270 | P = 0.117 | r = 0.441 | P = 0.008 |
| ML    | 0–2   | r = 0.431 | P = 0.001  | r = 0.184 | P = 0.182 | r = 0.515 | P = 0.000 |
|       | 2–3   | r = 0.639 | P = 0.000  | r = 0.519 | P = 0.001 | r = 0.516 | P = 0.001 |
| LPA   | 0–2   | r = 0.295 | P = 0.030  | r = 0.232 | P = 0.091 | r = 0.298 | P = 0.029 |
|       | 2–3   | r = 0.313 | P = 0.068  | r = 0.349 | P = 0.040 | r = 0.523 | P = 0.001 |
| Midvagina | 0–2   | r = 0.295 | P = 0.030  | r = 0.232 | P = 0.091 | r = 0.298 | P = 0.029 |
|       | 2–3   | r = 0.831 | P = 0.000  | r = 0.586 | P = 0.000 | r = 0.784 | P = 0.000 |
| Distal vagina | 0–2   | r = 0.394 | P = 0.019  | r = 0.295 | P = 0.030 | r = 0.232 | P = 0.091 |
|       | 2–3   | r = 0.394 | P = 0.019  | r = 0.295 | P = 0.030 | r = 0.232 | P = 0.091 |
| UH    | 0–2   | —       | —       | —       | —       | —       | —       |
|       | 2–3   | r = 0.708 | P = 0.000  | r = 0.784 | P = 0.000 | r = 0.784 | P = 0.000 |
| Mid-LA  | 0–2   | r = 0.120 | P = 0.491  | r = 0.207 | P = 0.232 | r = 0.479 | P = 0.004 |
|       | 2–3   | r = 0.120 | P = 0.491  | r = 0.207 | P = 0.232 | r = 0.479 | P = 0.004 |

LHL indicates length of LA; LHW, width of LA; midvagina, mean displacement of points 4 to 6.

In summary, quantified displacements on MRI could be used to hypothesize the mechanical mechanisms of PVP regarding the interaction between pressure and the support force contributing to the deformation of PVW and the supporting structures.

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