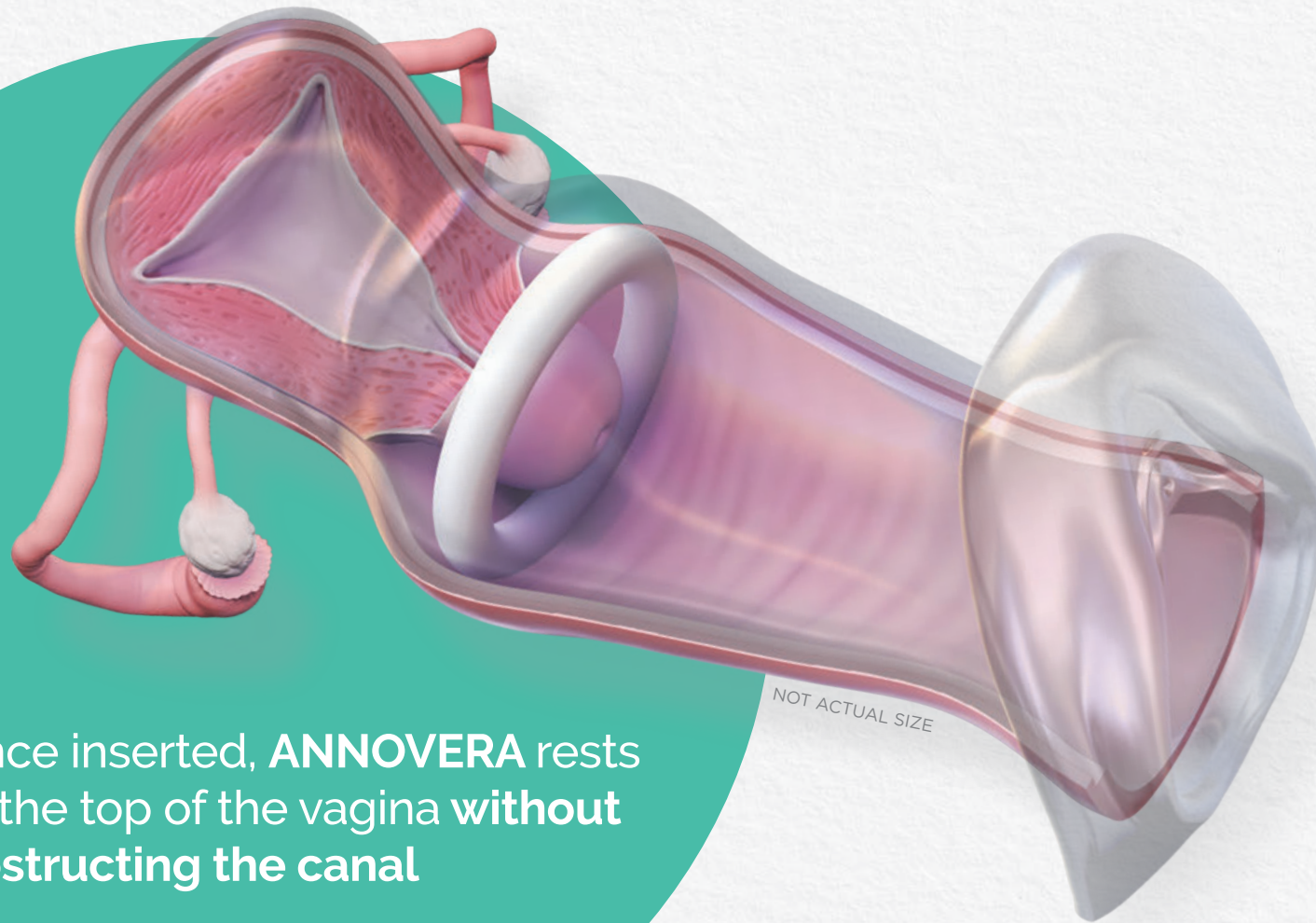
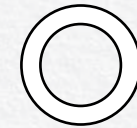


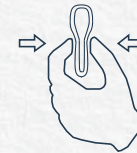
# The only long-lasting, reversible contraception that is patient-controlled and procedure-free



Once inserted, ANNOVERA rests at the top of the vagina without obstructing the canal



ANNOVERA is a **soft and comfortable** vaginal ring



Use thumb and index finger to **squeeze the ring into a narrow oval shape**



**Insert ANNOVERA** into your vagina **as far as possible**. [Click here](#) to download **Instructions for Use**



ANNOVERA is **inserted for 21 continuous days** and then **removed for 7 days** for 1 year (13 cycles)

## IMPORTANT SAFETY INFORMATION

**WARNING: CIGARETTE SMOKING AND SERIOUS CARDIOVASCULAR EVENTS**  
*See full prescribing information for complete boxed warning.*

- Females over 35 years old who smoke should not use ANNOVERA.
- Cigarette smoking increases the risk of serious cardiovascular events from combination hormonal contraceptive use.

Please see Indication and additional Important Safety Information on next page and Full Prescribing Information, including **BOXED WARNING** at [ANNOVERA.com/pi.pdf](http://ANNOVERA.com/pi.pdf)

**Annovera**<sup>®</sup>  
(segesterone acetate and ethinyl estradiol vaginal system)  
Delivers 0.15 mg/0.013 mg per day

## IMPORTANT SAFETY INFORMATION (CONT'D)

### CONTRAINDICATIONS

ANNOVERA is contraindicated and should not be used in women with a high risk of arterial or venous thrombotic diseases; current or history of breast cancer or other estrogen- or progestin-sensitive cancer; liver tumors, acute hepatitis, or severe (decompensated) cirrhosis; undiagnosed abnormal uterine bleeding; hypersensitivity to any of the components of ANNOVERA; and use of Hepatitis C drug combinations containing ombitasvir/paritaprevir/ritonavir, with or without dasabuvir.

### WARNINGS AND PRECAUTIONS

- Stop ANNOVERA if a thrombotic or thromboembolic event occurs, and at least 4 weeks before and through 2 weeks after major surgery. Start ANNOVERA no earlier than 4 weeks after delivery, in females who are not breastfeeding. Consider cardiovascular risk factors before initiating in all females, particularly those over 35 years.
- Discontinue if jaundice occurs.
- Stop ANNOVERA prior to starting therapy with the combination drug regimen ombitasvir/paritaprevir/ritonavir. ANNOVERA can be restarted 2 weeks following completion of this regimen.
- Do not prescribe ANNOVERA for females with uncontrolled hypertension or hypertension with vascular disease. Monitor blood pressure and stop use if blood pressure rises significantly in females with well-controlled hypertension.
- Monitor glucose in pre-diabetic or diabetic females taking ANNOVERA. Consider an alternate contraceptive method for females with uncontrolled dyslipidemias.
- Patients using ANNOVERA who have a significant change in headaches or irregular bleeding or amenorrhea should be evaluated. ANNOVERA should be discontinued if indicated.

- Other warnings include: gallbladder disease; depression; cervical cancer; increased serum concentrations of binding globulins; hereditary angioedema; chloasma (females who tend to develop chloasma should avoid exposure to the sun or UV radiation while using ANNOVERA); toxic shock syndrome (TSS) (if a patient exhibits symptoms of TSS, remove ANNOVERA, and initiate appropriate medical treatment); vaginal use (ANNOVERA may not be suitable for females with conditions that make the vagina more susceptible to vaginal irritation or ulceration).

### ADVERSE REACTIONS

The most common adverse reactions reported in at least 5% of women who received ANNOVERA were: headache/migraine, nausea/vomiting, vulvovaginal mycotic infection/candidiasis, lower/upper abdominal pain, dysmenorrhea, vaginal discharge, urinary tract infection, breast pain/tenderness/discomfort, bleeding irregularities including metrorrhagia, diarrhea, and genital pruritus.

### DRUG INTERACTIONS

Drugs or herbal products that induce certain enzymes, including CYP3A4, may decrease the effectiveness of ANNOVERA or increase breakthrough bleeding. Counsel patients to use a back-up or alternative method of contraception when enzyme inducers are used with ANNOVERA.

### INDICATION

ANNOVERA is a progestin/estrogen combination hormonal contraceptive indicated for use by females of reproductive potential to prevent pregnancy.

Limitations of Use: ANNOVERA has not been adequately studied in females with a body mass index  $>29$  kg/m<sup>2</sup>.

**Please note this information is not comprehensive. Please see Full Prescribing Information, including BOXED WARNING, at [ANNOVERA.com/pi.pdf](https://www.annovera.com/pi.pdf)**



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**Annovera**<sup>®</sup>  
(segesterone acetate and  
ethinyl estradiol vaginal system)  
Delivers 0.15 mg/0.013 mg per day