

Comments:	- Affix label here- Clinical Center/ID: ____ - ____ - ____ - ____ First Name _____ M.I. ____ Last Name _____
------------------	--

1. Date of Action: -- (M/D/Y)

₀ No

₁ Yes

2. Completed By: _____

3. Contact Type:

₁ Phone

₂ Mail

₃ Visit

₈ Other

4. Visit Type:

₁ Screening #

₂ Semi-Annual #

₃ Annual #

₄ Non-Routine

5. What study medication schedule did the participant follow?

HRT ___ ___ pills/week

CEE 0.3 mg ___ ___ pills/week

CEE 0.625 mg ___ ___ pills/week

MPA 2.5 mg ___ ___ pills/week

MPA 5 mg ___ ___ pills/week

MPA 10 mg ___ ___ pills/week

CaD ___ ___ pills/week

6. What is the new study medication schedule?
(Include all study medications the participant should take, including those that you are not changing.)

6.1. Medication:

6.2. Dosage:

1. ___ HRT: pills/week

2. ___ CEE 0.3 mg: pills/week

3. ___ CEE 0.625 mg: pills/week

4. ___ MPA 2.5 mg: pills/week

5. ___ MPA 5 mg: pills/week

6. ___ MPA 10 mg: pills/week

7. ___ CaD: pills/week

6.3 Is this a cyclic regimen?

K _____ V _____

7. Is the new study medication scheduled permanent?

- ₀ No →
- ₁ Yes

7.1. For how long should the participant follow this new study medication schedule?
(Record shortest length of time if more than one medication.)

weeks

8. Why did you make the change in the medication schedule?

8.1. HRT (Mark all that apply.)

- ₁ Bleeding
- ₂ Biopsy abnormality
- ₃ Abnormal transvaginal ultrasound
- ₄ Symptom intolerance
(Specify): _____

- ₈ Other
(Specify): _____

8.2. CaD (Mark all that apply.)

- ₁ Symptom intolerance
(Specify): _____

- ₈ Other
(Specify): _____
