

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Data Entry Form Version Log (01/06/2005)
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Form Name	Form Code	Version
Adverse Events and Serious Adverse Events	AE	V4.0.20050106
Adverse Events and Serious Adverse Events – Legend	AE-Legend	V2.0.20041004
NIH – Chronic Prostatitis Symptom Index	CPSI	V1.0.20040816
Demographics	DEMO	V1.0.20040816
Dispensing Log	DISP	V1.0.20040824
Drug Compliance	DCOMP	V1.0.20040816
Eligibility Checklist	ELIG	V1.0.20040816
EPS and Urine Testing	EUT	V1.0.20040824
Hospital Anxiety and Depression Scale©	HADS	V1.0.20040816
International Index of Erectile Function®	IIEF	V1.0.20040816
Male Sexual Health Questionnaire	MSHQ	V1.0.20040816
Medical History	MEDHX	V1.0.20040816
McGill Pain Questionnaire® (MPQ)	MCGILL	V1.0.20040816
Participant Expectations Questionnaire	EXP	V1.0.20050106
Physical Exam	EXAM	V1.0.20040816
Pre-screening Summary	PRESCR	V1.0.20040816
Randomization	RAND	V1.0.20040816
SF12 – Health Status Questionnaire®	SF12	V1.0.20040816
Study Stop Point	SSTOP	V1.0.20040816
Symptom Assessment	SYM	V1.0.20040816
Treatment Stop Point	TSTOP	V2.0.20041208
Unmasking Record	UNMASK	V2.0.20041004
Urine Screening	URINE	V1.0.20040816

Forms and Visit Schedule

Form Name	-1 to -4 Weeks (Screening)	0 Weeks (Randomization)	6 Weeks (Follow-up)	12 Weeks (Follow-up)
	Visit 1 (B1 Clinic)	Visit 2 (B2 Clinic)	Visit 3 (Clinic)	Visit 4 (Clinic)
Prescreening/Screening/Baseline				
Pre-Screening Summary (PRESCR)				
Informed Consent (Administrative)	X			
Medical History (MEDHX)	X			
Eligibility Checklist (ELIG)	X	X		
NIH-Chronic Prostatitis Symptom Index (CPSI)	X	X	X	X
Randomization (RAND)		X		
Procedures and Labs				
Adverse Events/Serious Adverse Events (AE)		X	X	X
Demographics (DEMO)	X			
Dispensing Log (DISP)		X	X	
Drug Compliance (DCOMP)			X	X
EPS and Urine Testing (EUT)	X			
Physical Exam (EXAM)	X			
Urine Screening (URINE)	X			
Symptom Questionnaires				
Symptom Assessment (SYM)	X	X	X (GRA)	X (GRA)
Health Status Questionnaire® (SF-12)		X	X	X
The McGill Pain Questionnaire® (MPQ)		X	X	X
Hospital Anxiety and Depression Scale© (HADS)		X	X	X
The International Index of Erectile Function® (IIEF)		X	X	X
The Male Sexual Health Questionnaire (MSHQ)		X	X	X
Participant Expectations Questionnaire (EXP)		X		
PRN Forms				
Study Stop Point (SSTOP)	PRN	PRN	PRN	X
Treatment Stop Point (TSTOP)	PRN	PRN	PRN	X
Unmasking Record (UNMASK)	PRN	PRN	PRN	PRN
Administrative Forms				
Clinical Center Staff “Signature and Delegation of Responsibilities” Log (STAFFLOG)	X	PRN	PRN	PRN
Concomitant Medication (CMED)	X	X	X	X
Participant ID Assignment Log (PTLOG)	X			
Participant Contact Information (PTCONT)	X			
Participant Transfer (TRANS)	PRN	PRN	PRN	PRN
Study Drug Tracking Log (TRACK)		X	X	X
Visit Checklist	X	X	X	X

Note: The two Eligibility forms are the only forms that must be entered for those participants who fail to be randomized.

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Adverse Events and Serious Adverse Events

To capture “pre-existing medical conditions” as defined in the MOP at Visit 2, please record any baseline signs and symptoms that the participant is experiencing. Beginning at Visit #3, list all adverse events (AEs) that have newly occurred, changed, been resolved, or are ongoing at EACH visit.

1. Were there any newly reported AEs; resolutions of AEs; follow-ups to ongoing AEs; or pre-existing medical conditions to be reported at this visit? ₁ Yes ₀ No

a. If YES, please list the total number of records at this visit: _____

Event Number	MedDRA Code	Date of Onset mm/dd/yyyy	Grade	Duration	Frequency	Relation ship to Study Drug	Action taken regarding Study Drug	Treatment for event	Outcome	Date of Resolution mm/dd/yyyy	Was the Event Serious?	Pre-existing condition
AE #	From CTC List	Use the Check Box if Event is continuous from previous visit.	record one	record one	record one	record one	record one	record one	record most appropriate	Use the Check Box if Event is continuous from this visit.		
_____	_____	____/____/____ <input type="checkbox"/>								____/____/____ <input type="checkbox"/>		
Specify Event (CTC Criteria):				Description of Event/Comments:								
_____	_____	____/____/____ <input type="checkbox"/>								____/____/____ <input type="checkbox"/>		
Specify Event (CTC Criteria):				Description of Event/Comments:								
_____	_____	____/____/____ <input type="checkbox"/>								____/____/____ <input type="checkbox"/>		
Specify Event (CTC Criteria):				Description of Event/Comments:								
_____	_____	____/____/____ <input type="checkbox"/>								____/____/____ <input type="checkbox"/>		
Specify Event (CTC Criteria):				Description of Event/Comments:								

Principal Investigator's Signature: _____ Date: ____/____/____ PI ID: _____
(MM/DD/YYYY)

**Randomized
Clinical Trial #1
(Protocol – CP01)**

Adverse Events and Serious Adverse Events

List all pre-existing medical conditions and/or adverse events that have newly occurred, changed, or resolved.

Grade	Duration	Frequency	Relationship to Study Drug	Action Taken Regarding Study Drug
1. Mild 2. Moderate 3. Severe 4. Life threatening or disabling 5. Fatal 88. Unknown 99. NA	1. Minutes 2. Hours 3. Days 88. Unknown 99. NA	1. Once 2. 2 to 3 episodes 3. 4 or more episodes 4. Daily 99. NA	1. Not related 2. Possibly related 3. Definitely related 88. Unknown/undetermined 99. NA	0. No action taken 1. Drug interrupted 2. Drug discontinued 99. N/A

Treatment for Event	Outcome	Was the Event Serious?	Pre-existing Condition
0. No 1. Yes 99. NA	1. Resolved/no follow-up needed 2. On-going/treatment continued *3. ER visit/prolonged hospitalization *4. Resulted in persistent or significant disability/incapacity *5. Congenital anomaly *6. Life threatening *7. Fatal 99. NA *Indicates Serious Adverse Events and must be reported to the IRB and DCC. PI signature needed on AEs and SAEs	0. No 1. Yes 99. NA	0. No 1. Yes 99. NA

C P C R N <i>Randomized Clinical Trial #1 (Protocol – CP01)</i>	Participant ID: _____	Participant Initials _____
	Clinical Center: _____	Visit Number _____
	CRF Date: ____/____/____	RC ID: _____

NIH – CHRONIC PROSTATITIS SYMPTOM INDEX
PARTICIPANT COMPLETES AT VISITS 1, 2, 3, AND 4.

Pain or Discomfort

1. In the last week, have you experienced any pain or discomfort in the following areas?
 - a. Area between rectum and testicles (perineum) ₁ Yes ₀ No
 - b. Testicles ₁ Yes ₀ No
 - c. Tip of the penis (not related to urination) ₁ Yes ₀ No
 - d. Below your waist, in you pubic or bladder area ₁ Yes ₀ No

2. In the last week, have you experienced:
 - a. Pain or burning during urination? ₁ Yes ₀ No
 - b. Pain or discomfort during or after sexual climax (ejaculation)? ₁ Yes ₀ No

3. How often have you had pain or discomfort in any of these areas over the last week?
 - ₀ Never
 - ₁ Rarely
 - ₂ Sometimes
 - ₃ Often
 - ₄ Usually
 - ₅ Always

4. Which number best describes your AVERAGE pain or discomfort on the days that you had it, over the last week?

<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
0	1	2	3	4	5	6	7	8	9	10
No Pain							Pain as bad as you can imagine			

Urination

5. How often have you had a sensation of not emptying your bladder completely after you finished urinating, over the last week?
 - ₀ Not at all
 - ₁ Less than 1 time in 5
 - ₂ Less than half the time
 - ₃ About half the time
 - ₄ More than half the time
 - ₅ Almost always

6. How often have you had to urinate again less than two hours after you finished urinating, over the last week?
 - ₀ Not at all
 - ₁ Less than 1 time in 5
 - ₂ Less than half the time
 - ₃ About half the time
 - ₄ More than half the time
 - ₅ Almost always

C P C R N <i>Randomized Clinical Trial #1 (Protocol – CP01)</i>	Participant ID: _____	Participant Initials _____
	Clinical Center: _____	Visit Number _____
	CRF Date: ____/____/____	RC ID: _____

NIH – CHRONIC PROSTATITIS SYMPTOM INDEX
PARTICIPANT COMPLETES AT VISITS 1, 2, 3, AND 4.

Impact of Symptoms

7. How much have your symptoms kept you from doing the kinds of things you would usually do, over the last week?
- ₀ None
₁ Only a little
₂ Some
₃ A lot
8. How much did you think about your symptoms, over the last week?
- ₀ None
₁ Only a little
₂ Some
₃ A lot

Quality of Life

9. If you were to spend the rest of your life with your symptoms just the way they have been during the last week, how would you feel about that?
- ₀ Delighted
₁ Pleased
₂ Mostly satisfied
₃ Mixed (about equally satisfied and dissatisfied)
₄ Mostly dissatisfied
₅ Unhappy
₆ Terrible

To be completed by the RC:

Overall Score: _____
(Sum of # 1 – 9)

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/_____	RC ID: _____

Study Drug Compliance

RESEARCH COORDINATOR TO COMPLETE AT VISITS 3 AND 4.

1. Treatment Point: ₁ 6 Weeks ₂ 12 Weeks

2. **Study Drug Returned:**

Amount Dispensed (A)	Amount Returned (B)	Amount Lost/Destroyed (C)	Amount Used A-(B+C) (D)	Amount that should have been used (E)	Percent Compliance (D/E)X100 (F)

Date (Month/Day/Year) (G)	Initials (H)

3. ***If the participant is less than 80% compliant, please provide reason(s):***

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Demographics

PARTICIPANT COMPLETES AT VISIT 1.

1. What is your date of birth? _____ / _____ / _____ (MM/DD/YYYY)
2. What is your gender? ₁ Male ₂ Female
3. What do you consider to be your ethnicity? ₁ Hispanic or Latino ₂ Not Hispanic or Latino
4. Using the categories below, what do you consider to be your racial background?
 - a. North American Indian/Northern Native ₁ Yes ₀ No
 - b. Asian/Asian American ₁ Yes ₀ No
 - c. Black/African American ₁ Yes ₀ No
 - d. Native Hawaiian/Other Pacific Islander ₁ Yes ₀ No
 - e. White/Caucasian ₁ Yes ₀ No
 - f. Other (Please specify) _____ ₁ Yes ₀ No
5. What is the highest educational level you have attained?
 - ₁ Less than high school
 - ₂ High school or GED
 - ₃ Some college
 - ₄ Graduated from college/university
 - ₅ Graduate or professional school after college/university
6. What is your current employment status?
 - ₁ Employed
 - ₂ Unemployed
 - ₃ Retired
 - ₄ Full-time homemaker
 - ₅ Disabled
7. What is your annual family income? (In U.S. dollars, please see the MOP for conversion method)
 - ₁ \$10,000 or less
 - ₂ \$10,001 to \$25,000
 - ₃ \$25,001 to \$50,000
 - ₄ \$50,001 to \$100,000
 - ₅ More than \$100,000
 - ₉₉ Prefer not to Answer
8. What is your ZIP (US) or Postal (Canadian) Code?
 - a. ZIP Code (for US Residents) _____ N/A
 - b. Postal Code (for Canadian Residents) _____ N/A
9. Have any family members ever been diagnosed with Painful Bladder Syndrome (PBS) / Interstitial Cystitis (IC)? ₁ Yes ₀ No ₈₈ Unknown
10. Have any family members ever been diagnosed with Chronic Pelvic Pain Syndrome (CPPS) / Chronic Prostatitis (CP)? ₁ Yes ₀ No ₈₈ Unknown
11. Are you living with a spouse or partner? ₁ Yes ₀ No

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Dispensing Log

RESEARCH COORDINATOR COMPLETES AT VISITS 2 AND 3.

1. Date study drug capsules/tablets dispensed: _____/_____/_____

2. Total number of study drug capsules/tablets dispensed: _____
(Record on Study Medication Tracking Log)

Affix and sign the drug label for the study drug capsules/tablets here.
Record on Study Medication Tracking Log

NOTE: By signing the label here, you are confirming that you have:

1. Checked the label on the bottle with the randomization number on the Randomization form, and
2. Confirmed that the drug is being given to the participant with the ID number written on this form.

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Eligibility Checklist

RESEARCH COORDINATOR COMPLETES AT VISITS 1 AND 2.

Inclusion Criteria

Complete questions #1, #2 and #3 at Visit #1 only:

- | | |
|--|--|
| 1. Participant has signed and dated the appropriate Informed consent document. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| a. If Yes , record date the form was signed | ____/____/____ |
| 2. Participant is male. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 3. Participant is ≥ 18 years of age. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 4. Participant has an overall score on the NIH-CPSI of ≥ 12 out of a potential of 0-43 points. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| a. Record the overall NIH-CPSI score.
<i>(Please refer to the overall score on the CPSI form)</i> | _____ |
| 5. Participant has had symptoms of discomfort or pain in the pelvic region for at least a six-week interval at the time of presentation. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 6. Symptoms bothersome enough to prompt a physician visit have been present for two years or less. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |

ALL INCLUSION CRITERIA RESPONSES MUST BE “YES” FOR THE PARTICIPANT TO BE ELIGIBLE FOR ENROLLMENT

Exclusion Criteria

Complete question #7 at Visit #2 only:

- | | |
|---|--|
| 7. Participant has evidence of facultative Gram negative or enterococcus with a value of ≥ 1,000 CFU/ml in mid-stream urine (VB2). | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| a. Record the actual urine count in CFU/ml: | _____ |
| 8. Participant has previously received alfuzosin (Uroxatral®), tamsulosin hydrochloride (Flomax®), doxazosin mesylate (Cardura®), terazosin HCL (Hytrin®), or other alpha-adrenergic receptor blockers for symptoms of CP/CPPS or within the past two years for any other reason. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 9. Participant has a history of prostate, penile, testicular, bladder, or urethral cancer or has undergone pelvic radiation, systemic chemotherapy, or intravesical chemotherapy. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 10. Participant has a history of moderate or severe hepatic impairment, severe renal insufficiency, severe or unstable cardiovascular (i.e. prolonged QT), respiratory, hematological, endocrinological, neurological, or other somatic disorders. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 11. Participant has unilateral orchialgia without pelvic symptoms, active urethral stricture, or neurological disease or disorder affecting the bladder. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 12. Participant has uninvestigated, significant hematuria. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Eligibility Checklist

RESEARCH COORDINATOR COMPLETES AT VISITS 1 AND 2.

- | | |
|--|--|
| 13. Participant has undergone TURP, TUIP, TUIBN, TUMT, TUNA, balloon dilation of the prostate, open prostatectomy or any other prostate surgery or treatment such as cryotherapy or thermal therapy. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 14. Participant has a neurological impairment or psychiatric disorder preventing his understanding of consent and his ability to comply with the protocol. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 15. Participant is currently taking exclusionary medications such as potent CYP3A4 inhibitors (i.e. ketoconazole, itraconazole, or ritonavir) or erythromycin. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |

ALL EXCLUSION CRITERIA RESPONSES MUST BE “NO” FOR THE PARTICIPANT TO BE ELIGIBLE FOR ENROLLMENT

Deferral Criteria

- | | |
|---|--|
| 16. Participant has had a urinary tract infection, with a urine culture value of >100,000 CFU/ml, in the past three (3) months. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 17. Participant has had clinical evidence of urethritis, i.e. including urethral discharge or positive culture, in the past three (3) months, diagnostic of the following sexually transmitted diseases (STDs): gonorrhea, chlamydia, mycoplasma, or trichomonas. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 18. Participant has had a prostate biopsy in the past three (3) months. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 19. Participant has experienced symptoms of acute or chronic epididymitis in the past three (3) months. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 20. Participant has been diagnosed with or treated for symptomatic genital herpes in the past twelve (12) months. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 21. Participant has been taking excluded medications such as Cialis®, Levitra® and Viagra® in the past one (1) week. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |
| 22. Participant has been taking prescription drugs with 5-alpha reductase activity (i.e. dutasterade or finasteride) in the past twelve (12) months. | <input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No |

23. Did the participant meet all eligibility criteria at this visit?	<input type="checkbox"/> ₁ Yes <input type="checkbox"/> ₀ No
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C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/_____	RC ID: _____

EPS and Urine Testing

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

1. Date of participant's EPS and Urine Test _____ (MM/DD/YYYY)
2. Has the participant remained abstinent for the past 48 hours? ₁ Yes ₀ No

The table below lists the species to be identified in each sample, and each species's appropriate code. Use these codes when completing the tables for the culture count for each modified four glass test sample.

Species	Species Code
Enterococcus Fecalis	07
Escherichia Coli	09
Klebsiella	10
Pseudomonas	11
Proteus	12
Other Enterobacteracea	13

VB2 - 48 Hour Culture

3. Date of 48 hour culture _____ (MM/DD/YYYY)
4. Was there any growth? ₁ Yes ₀ No

If **Yes**, please complete the chart below, indicating what species were present, and the culture count measured in CFU/ml:

Enter total number of species recorded: _____

Species Code	Please enter actual count
_____	_____,_____ CFU/ml
_____	_____,_____ CFU/ml
_____	_____,_____ CFU/ml
_____	_____,_____ CFU/ml
_____	_____,_____ CFU/ml
_____	_____,_____ CFU/ml

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
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		CRF Date: ____/____/____	RC ID: _____

EPS and Urine Testing
RESEARCH COORDINATOR COMPLETES AT VISIT 1.

EPS

5. Was the participant able to provide an EPS sample? ₁ Yes ₀ No

If **No**, please go to question #8.

If **Yes**, please complete questions #6 and #7.

6. Estimated volume of EPS sample ₁ None
₂ 1 or 2 drops
₃ 3 or more drops

7. White Blood Cell Count (/hpf) ₁ ≤ 25
₂ 26 - 50
₃ 51-75
₄ 76-100
₅ > 100

a. If ≤ 25, give actual count _____ /hpf

VB3

8. Was the participant able to provide a VB3 sample? ₁ Yes ₀ No

If **Yes**, please complete questions #9.

9. White Blood Cell Count (/hpf) ₁ ≤ 25
₂ 26 - 50
₃ 51-75
₄ 76-100
₅ > 100

a. If ≤ 25, give actual count _____ /hpf

C P R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Physical Exam

Clinician completes at Visits 1.

1. Height:
 - a. Feet _____
 - b. Inches _____
2. Weight: _____ lbs.
3. Blood Pressure:
 - a. Systolic (mmHg) _____
 - b. Diastolic (mmHg) _____
4. Abdominal exam: ₁ Normal ₀ Abnormal

Pelvic Exam:

5. External Genitalia: ₁ Normal ₀ Abnormal ₉₉ Not Applicable
 - a. If **Abnormal** please specify: _____
6. Rectal / Bimanual exam: ₁ Normal ₀ Abnormal ₉₉ Not Applicable

Men only (Check N/A for women)

7. Suprapubic Tenderness ₁ Yes ₀ No ₉₉ Not Applicable
8. Penis Circumcised ₁ Yes ₀ No ₉₉ Not Applicable
9. Prostate
 - a. Enlarged ₁ Yes ₀ No ₉₉ Not Applicable
 - b. Irregular ₁ Yes ₀ No ₉₉ Not Applicable
 - c. Tender ₁ Yes ₀ No ₉₉ Not Applicable
 - d. Pelvic Wall (muscle) Tenderness ₁ Yes ₀ No ₉₉ Not Applicable

Women only (Check N/A for males & post-menopausal women)

10. Pelvic floor musculature tenderness ₁ Yes ₀ No ₉₉ Not Applicable
11. Uterus
 - a. Absent ₁ Yes ₀ No ₉₉ Not Applicable
 - b. Normal ₁ Yes ₀ No ₉₉ Not Applicable
 - c. Abnormal ₁ Yes ₀ No ₉₉ Not Applicable
If **Yes** please specify: _____
12. Pelvic organ support
 - a. Normal ₁ Yes ₀ No ₉₉ Not Applicable
 - b. Prolapse present, no vaginal points beyond the hymen ₁ Yes ₀ No ₉₉ Not Applicable
 - c. Prolapse present, at least one vaginal point beyond the hymen ₁ Yes ₀ No ₉₉ Not Applicable
13. Menstruating ₁ Yes ₀ No ₉₉ Not Applicable
 - a. If **Yes**, date of last menstrual period: Date: ____/____/____ (MM/DD/YYYY)
 - b. If **No**: ₁ Premenopausal
₂ Prior Hysterectomy
₃ Postmenopausal

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
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		CRF Date: ____/____/____	RC ID: _____

Participant Expectations Questionnaire

PARTICIPANT COMPLETES AT VISIT 2.

As a participant in this study, we are very interested in what you expect.

1. Do you expect that participating in this study will help your pelvic pain/prostatitis symptoms?

₁ Yes
 ₀ No
 ₈₈ Unknown

2. Based on what you have heard about the study, how much do you expect your symptoms to change by the end of this study?

₁ Markedly worse
₂ Moderately worse
₃ Slightly worse
₄ No change
₅ Slightly improved
₆ Moderately improved
₇ Markedly improved

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
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		CRF Date: ____/____/____	RC ID: _____

Medical History

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

I'm going to ask you some questions . . .

1. Have you ever been diagnosed with interstitial cystitis ("IC"), chronic prostatitis, or chronic pelvic pain symptoms? ₁ Yes ₀ No
 - a. If **YES**, at what age were you diagnosed? _____ age
2. Do you know when your IC, chronic prostatitis, or chronic pelvic pain symptoms first began? ₁ Yes ₀ No
 - a. If **YES**, at what age did they first begin? _____ age

I am going to ask you some questions about some medical disorders and conditions. Please tell me if you have ever been diagnosed with any of the following:

Genitourinary Disorders: (Both Men and Women)

3. Bladder cancer ₁ Yes ₀ No ₈₈ U/K
4. Urinary Tract Infection ₁ Yes ₀ No ₈₈ U/K

(Women only)

5. Pelvic Inflammatory Disease (PID) ₁ Yes ₀ No ₈₈ U/K
6. Endometriosis ₁ Yes ₀ No ₈₈ U/K
7. Vulvodynia ₁ Yes ₀ No ₈₈ U/K
8. Gynecologic cancer ₁ Yes ₀ No ₈₈ U/K
9. Vulvovestibulitis ₁ Yes ₀ No ₈₈ U/K

(Men only)

10. Acute / Chronic Bacterial prostatitis ₁ Yes ₀ No ₈₈ U/K
11. Epididymitis ₁ Yes ₀ No ₈₈ U/K

Respiratory Tract Disorders/Allergies: (Both Men and Women)

12. Have you been diagnosed with having any respiratory tract disorders and/or allergies? ₁ Yes ₀ No ₈₈ U/K

If **Yes**, which of the following:

 - a. Asthma ₁ Yes ₀ No ₈₈ U/K
 - b. Drug allergies ₁ Yes ₀ No ₈₈ U/K
 - c. Food allergies ₁ Yes ₀ No ₈₈ U/K
 - d. Skin allergies (contact dermatitis) ₁ Yes ₀ No ₈₈ U/K

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Medical History

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

- | | | | |
|--------------------------------|---|--|--|
| e. Sinusitis | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| f. Hayfever, allergic rhinitis | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| g. Latex allergies | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| h. Other allergies | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Gastrointestinal Disease (Both Men and Women)

- | | | | |
|--|---|--|--|
| 13. Have you been diagnosed with having any gastrointestinal diseases? | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| If Yes , which of the following: | | | |
| a. Irritable bowel syndrome | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| b. Diverticulitis | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. Ulcerative Colitis/Crohn's Disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Other gastrointestinal disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Endocrine or metabolic disease (Both Men and Women)

- | | | | |
|--|---|--|--|
| 14. Have you been diagnosed with having any endocrine or metabolic diseases? | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| If Yes , which of the following: | | | |
| a. Diabetes | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| b. Hypothyroid disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. Hyperthyroid disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Other endocrine or metabolic disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Hematopoietic, lymphatic, or infectious disease (Both Men and Women)

- | | | | |
|---|---|--|--|
| 15. Have you been diagnosed with having any blood, lymphatic, or infectious diseases? | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| If Yes , which of the following: | | | |
| a. Epstein-Barr virus/Chronic Fatigue Syndrome | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| b. Tuberculosis | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. HIV/AIDS | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Viral Hepatitis (A,B,C,D,E) | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Medical History

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

Psychiatric Disease (Both Men and Women)

16. Have you been diagnosed with having any psychiatric diseases? ₁ Yes ₀ No ₈₈ U/K
- If **Yes**, which of the following:
- a. Depression ₁ Yes ₀ No ₈₈ U/K
 - b. Eating disorder ₁ Yes ₀ No ₈₈ U/K
 - c. Anxiety/panic attacks ₁ Yes ₀ No ₈₈ U/K
 - d. Suicide attempt ₁ Yes ₀ No ₈₈ U/K
 - e. Other psychiatric disease ₁ Yes ₀ No ₈₈ U/K

Sexually Transmitted Disease (Both Men and Women)

17. Have you been diagnosed with having any sexually transmitted diseases? ₁ Yes ₀ No ₈₈ U/K
- If **Yes**, which of the following:
- a. Gonorrhea ₁ Yes ₀ No ₈₈ U/K
 - b. Syphilis ₁ Yes ₀ No ₈₈ U/K
 - c. Chlamydia ₁ Yes ₀ No ₈₈ U/K
 - d. HIV/AIDS ₁ Yes ₀ No ₈₈ U/K
 - e. Genital herpes ₁ Yes ₀ No ₈₈ U/K
 - f. Genital warts ₁ Yes ₀ No ₈₈ U/K
 - g. Trichomonas ₁ Yes ₀ No ₈₈ U/K
 - h. Other sexually transmitted disease ₁ Yes ₀ No ₈₈ U/K

(Men only)

If **Yes**, please respond to the following:

- i. Nongonococcal Urethritis ₁ Yes ₀ No ₈₈ U/K

Cardiovascular Disease (Both Men and Women)

18. Have you been diagnosed with having any cardiovascular diseases? ₁ Yes ₀ No ₈₈ U/K
- If **Yes**, which of the following:
- a. Hypertension ₁ Yes ₀ No ₈₈ U/K

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Medical History

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

- | | | | |
|---|---|--|--|
| b. High cholesterol | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. Coronary artery disease (heart attack, chest pain) | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Stroke | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| e. Arrhythmia | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Neurologic Disease (Both Men and Women)

19. Have you been diagnosed with having any neurological diseases? ₁ Yes ₀ No ₈₈ U/K

If **Yes**, which of the following:

- | | | | |
|---------------------------------------|---|--|--|
| a. Lumbosacral/Vertebral Disc Disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| b. Numbness or tingling in limbs | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. History of seizuers | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Migraine headaches | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| e. Peripheral Neuropathy | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| f. Other neurological disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Autoimmune/Other Disorders: (Both Men and Women)

20. Have you been diagnosed with having any autoimmune/ other disorders? ₁ Yes ₀ No ₈₈ U/K

If **Yes**, which of the following:

- | | | | |
|---|---|--|--|
| a. Fibromyalgia or Fibromyositis | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| b. Autommune Disorders (ex. Lupus, Rheumatoid Arthritis, Sjogren’s Scleroderma) | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. Other musculoskeletal, rheumatologic, or connective tissue disease | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

Now I am going to ask some questions about some surgeries that you may have had.

(Women Only)

Urological/Gynecologic Surgeries:

21. Have you ever had urological/gynecologic surgeries? ₁ Yes ₀ No ₈₈ U/K

If **Yes**, please respond to the following:

- | | | | |
|---------------------------------|---|--|--|
| a. Pelvic organ prolapse repair | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
|---------------------------------|---|--|--|

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
	Clinical Center: _____	Visit Number _____	
	CRF Date: ____/____/____	RC ID: _____	

Medical History

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

- | | | | |
|-------------------------|---|--|--|
| b. Hysterectomy | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| c. Oophorectomy | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |
| d. Incontinence surgery | <input type="checkbox"/> ₁ Yes | <input type="checkbox"/> ₀ No | <input type="checkbox"/> ₈₈ U/K |

22. How many children have you given birth to by the following:

- | | | | |
|-------------------------|-------|---|--|
| a. By vaginal delivery | _____ | <input type="checkbox"/> Not Applicable | |
| b. By caesarean section | _____ | <input type="checkbox"/> Not Applicable | |

23. Are you postmenopausal? ₁ Yes ₀ No ₈₈ U/K

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

The Male Sexual Health Questionnaire

PARTICIPANT COMPLETES AT VISIT 2, 3, AND 4.

The following questions deal with male ejaculation and the pleasure you have with ejaculation. Ejaculation or “cumming” is the release of semen or “cum” during sexual climax. These questions concern all of your ejaculations when having sexual activity. These could include ejaculations you have had with your main partner, as well as with other partners, or ejaculations you have had when masturbating. Please **check only one response** for each question.

1. **In the last month**, how often have you been able to ejaculate when having sexual activity?
 - ₅ All of the time
 - ₄ Most of the time
 - ₃ About half of the time
 - ₂ Less than half of the time
 - ₁ None of the time/Could not ejaculate

2. **In the last month**, when having sexual activity, how often did you feel that you took too long to ejaculate or “cum”?
 - ₅ None of the time
 - ₄ Less than half of the time
 - ₃ About half of the time
 - ₂ Most of the time
 - ₁ All of the time
 - ₀ Could not ejaculate

3. **In the last month**, when having sexual activity, how often have you felt like you were ejaculating (“cumming”), but no fluid came out?
 - ₅ None of the time
 - ₄ Less than half of the time
 - ₃ About half of the time
 - ₂ Most of the time
 - ₁ All of the time
 - ₀ Could not ejaculate

4. **In the last month**, how would you rate the strength or force of your ejaculation?
 - ₅ As strong as it always was
 - ₄ A little less strong than it used to be
 - ₃ Somewhat less strong than it used to be
 - ₂ Much less strong than it used to be
 - ₁ Very much less strong than it used to be
 - ₀ Could not ejaculate

5. **In the last month**, how would you rate the amount or volume of semen when you ejaculate?
 - ₅ As much as it always was
 - ₄ A little less than it used to be
 - ₃ Somewhat less than it used to be
 - ₂ Much less than it used to be
 - ₁ Very much less than it used to be
 - ₀ Could not ejaculate

6. **Compared to ONE month ago**, would you say the physical pleasure you feel when you ejaculate has...
 - ₅ Increased a lot
 - ₄ Increased moderately
 - ₃ Neither increased or decreased
 - ₂ Decreased moderately
 - ₁ Decreased a lot
 - ₀ Could not ejaculate

7. **In the last month**, have you experienced any physical pain or discomfort when you ejaculated? Would you say you have...
 - ₅ No pain at all
 - ₄ Slight amount of pain or discomfort
 - ₃ Moderate amount of pain or discomfort
 - ₂ Strong amount of pain or discomfort
 - ₁ Extreme amount of pain or discomfort
 - ₀ Could not ejaculate

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

The Male Sexual Health Questionnaire

PARTICIPANT COMPLETES AT VISIT 2, 3, AND 4.

8. **In the last month**, if you have had any ejaculation difficulties or have been unable to ejaculate, have you been bothered by this?
- ₅ Not at all bothered
 - ₄ A little bit bothered
 - ₃ Moderately bothered
 - ₂ Very bothered
 - ₁ Extremely bothered

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Randomization

RESEARCH COORDINATOR COMPLETES AT VISIT 2.

1. Did the participant meet all of the eligibility criteria at the first baseline visit? ₁ Yes ₀ No
2. Does the participant still meet all of the eligibility criteria at the second baseline visit? ₁ Yes ₀ No
3. Record randomization number _____

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Study Stop Point

RESEARCH COORDINATOR COMPLETES AT VISIT 4, OR IF THE PARTICIPANT WITHDRAWS FROM STUDY PRIOR TO VISIT 4.

1. Has the participant successfully completed this study? ₁ Yes ₀ No

If **No**, indicate reason for withdrawal:

- a. No longer willing to follow the protocol/interested in participating ₁ Yes ₀ No
- b. Participant dissatisfied with treatment ₁ Yes ₀ No
- c. Lost to follow-up ₁ Yes ₀ No
- d. Participant has personal constraints ₁ Yes ₀ No
- e. Adverse Event/Serious Adverse Event ₁ Yes ₀ No
- f. Physician's Discretion ₁ Yes ₀ No
- g. Other (Specify: _____) ₁ Yes ₀ No

2. Visit Number that the participant was last seen: _____

3. Date that the participant was last seen: _____
(MM/DD/YYYY)

The following questions are for Participant Close-Out. (Participant and Research Coordinator complete at final study visit or premature termination from the study.)

4. Do you think the current status of your symptoms is related to the study medications? ₁ Yes ₀ No

5. Which medication do you think you received?
₁ Alfuzosin
₂ Placebo
₈₈ Couldn't tell

6. Referring to your response in question #5, what made you think that?
₁ CP was better
₂ CP was worse
₃ CP remained unchanged
₄ Experienced side effects
₅ Did not experience side effects
₉₈ Other: _____

Research Coordinator completes question #7:

7. Which medication do you think the participant received?
₁ Alfuzosin
₂ Placebo
₈₈ Couldn't tell

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Study Stop Point

RESEARCH COORDINATOR COMPLETES AT VISIT 4, OR IF THE PARTICIPANT WITHDRAWS FROM STUDY PRIOR TO VISIT 4.

The following section is for Study Close-out.

(PRINCIPAL INVESTIGATOR AND RESEARCH COORDINATOR COMPLETE WHEN PARTICIPANT STOPS PARTICIPATION IN THE STUDY.)

8. Physician Comments (optional): _____

SIGNATURES: Please complete the following section regardless of the reason for termination of study participation.

I verify that all information collected on the CPCRN-2 data collection forms for this participant is correct to the best of my knowledge and was collected in accordance with the procedures outlined in the CPCRN-2 Protocol and Manual of Procedures.

 Principal Investigator's Signature Date: ____/____/____
(MM/DD/YYYY)

9. Did the PI sign this form? Yes No

 Research Coordinator's Signature Date: ____/____/____
(MM/DD/YYYY)

10. Did the RC sign this form? Yes No

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Symptom Assessment

PARTICIPANT COMPLETES AT VISITS 1, 2, 3 AND 4.

1. Think about the pain/discomfort associated with your bladder, prostate, and/or pelvic region.
 - a. **On average**, how would you rate this **pain/discomfort** during the past 4 weeks?
(Please **circle** the number that best describes this **pain/discomfort**.)

No pain/discomfort											Most severe pain I can imagine
0	1	2	3	4	5	6	7	8	9	10	
 - b. How long has this pain/discomfort described in Question #1a been present?
(Please **check** the option that best describes your answer.)

₁ less than 4 weeks ₂ 4 weeks or more

2. Urgency is defined as the urge or pressure to urinate.
 - a. **On average**, how would you rate the **urgency** that you have felt during the past 4 weeks?
(Please **circle** the number that best describes this **urgency**.)

No urgency											Most severe urgency I can imagine
0	1	2	3	4	5	6	7	8	9	10	
 - b. How long has this urgency described in Question #2a been present?
(Please **check** the option that best describes your answer.)

₁ less than 4 weeks ₂ 4 weeks or more

3. Think about your frequency of urination compared to what you consider to be normal.
 - a. **On average**, how would you rate your frequency of urination during the past 4 weeks?
(Please **circle** the number that best describes this **frequency**.)

Totally normal											Most severe frequency I can imagine
0	1	2	3	4	5	6	7	8	9	10	
 - b. How long has this frequency described in Question #3a been present?
(Please **check** the option that best describes your answer.)

₁ less than 4 weeks ₂ 4 weeks or more
 - c. **On average**, during the past 4 weeks, how many times did you urinate in a 24-hour period?
(Please **check** the option that best describes your answer.)

₁ 6 times or less ₂ 7-10 times ₃ 11 – 14 times ₄ 15 times or more

(Question #4 is for treatment/follow-up visits only)

4. As compared to when you started the study, how would you rate your overall symptoms now?

₁ Markedly worse
₂ Moderately worse
₃ Slightly worse
₄ No change
₅ Slightly improved
₆ Moderately improved
₇ Markedly improved

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Treatment Stop Point

RESEARCH COORDINATOR COMPLETES AT VISIT 4, OR IF THE PARTICIPANT WITHDRAWS FROM TREATMENT PRIOR TO VISIT 4.

1. Date participant took final dose of study agent. ____/____/____
(MM/DD/YYYY)

2. Has the participant stopped treatment early? ₁ Yes ₀ No

3. Use of unacceptable medication ₁ Yes ₀ No

4. Adverse Event, *as determined by the PI* ₁ Yes ₀ No
 - a. Listed on Adverse Event form as AE #: _____
 - b. Date of onset: ____/____/____
(MM/DD/YYYY)
 - c. Please specify: _____

5. Adverse Event, *as determined by the Participant* ₁ Yes ₀ No
 - a. Listed on Adverse Event form as AE #: _____
 - b. Date of onset: ____/____/____
(MM/DD/YYYY)
 - c. Please specify: _____

6. Participant dissatisfied with treatment ₁ Yes ₀ No
 - a. Please specify reason: _____

7. Participant no longer interested in participating ₁ Yes ₀ No
 - a. Please specify reason: _____

8. Participant was lost to follow up. ₁ Yes ₀ No
 - a. Please specify reason: _____

9. Participant has personal constraints ₁ Yes ₀ No
 - a. Please specify reason: _____

10. Physician's Discretion ₁ Yes ₀ No
 - a. Please specify reason: _____

11. Other ₁ Yes ₀ No
 - a. Please specify reason: _____

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Unmasking Record

Research Coordinator completes if the study drug needs to be unmasked.

Photocopies of this form with signatures must be sent to the DCC.

1. Date of unmasking: _____ / _____ / _____
(MM/DD/YYYY)
2. Time of unmasking: _____ : _____ ₁ AM ₂ PM
3. Randomization Number: _____
4. Was the DCC contacted within 24 hours of unmasking? ₁ Yes ₀ No
 - a. If **Yes**, name of person contacted: _____
 - b. If **No**, state the reason: _____
5. Was the study drug unmasked because of an SAE? ₁ Yes ₀ No
 - a. Listed on Adverse Event form as AE # _____

P.I. Signature: _____ Date: ____/____/____ PI ID: _____
(MM/DD/YYYY)

Directions: Fax this form to the DCC at (215) 573-4790.

C P C R N 2	Randomized Clinical Trial #1 (Protocol – CP01)	Participant ID: _____	Participant Initials _____
		Clinical Center: _____	Visit Number _____
		CRF Date: ____/____/____	RC ID: _____

Urine Screening

RESEARCH COORDINATOR COMPLETES AT VISIT 1.

1. Date urine sample obtained: _____ / _____ / _____ (MM/DD/YYYY)

2. Dipstick Urinalysis: ₁ Normal ₀ Abnormal

If **ABNORMAL**, please check if present:

a. Nitrite ₁ Yes ₀ No

b. Occult Blood ₁ Yes ₀ No

c. Leukocytes ₁ Yes ₀ No

d. Protein / Ketones ₁ Yes ₀ No

3. Did this participant have a positive urine culture (colony count of more **10³** of uropathogens)? ₁ Yes ₀ No

(Question #s 4 and 5 for males only, check N/A for females)

4. Date residual urine volume measured: _____ / _____ / _____ (MM/DD/YYYY)

OR N/A

5. Was the residual urine volume greater than 100 cc as measured by ultrasound or catheter? ₁ Yes ₀ No ₉₉ N/A