

Patient number

Date of report
month day year

NIH - BPH TRIAL

ADVERSE EVENT REPORT

This form is to be completed if the patient has had any adverse experiences, drug reactions, side effects, abnormal laboratory values, hospitalizations, discontinued coded medications, other complications or pre-existing conditions that worsened.

If this is a **SERIOUS** event or the patient discontinued coded medications, this form should be **FAXED** to the Biostatistical Coordinating Center **IMMEDIATELY** at (301) 881-3589.

A. Report Identification

1. Clinic number (CLINIC)

2. Patient Identification number (Complete a **OR** b)

a. If before randomization, Screening number (SCREEN)

S

b. If after randomization, Patient number (PATID)

<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>
clinic			patient		

3. Patient's initials (INITS)

<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="text"/>
first		last	

4. Patient's date of birth (DOB)

<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>
month	day	year

5. Date of report (ZRIRD)

<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>
month	day	year

6. Date of onset of adverse experience (ZRIODT)

<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>	<input type="text"/> <input type="text"/>
month	day	year

7. Type of report (ZRITYP)

☐ Initial

☐ Follow-up

B. General Classification

1. Adverse experience (short description) (ZGICAE) (ZGCOSTG)

2. Did the adverse experience result in:
(Check all that apply)

☐ Death (ZGCDEA)

☐ Required or prolonged hospitalization (ZGCHOSP)

☐ Permanent or severe disability (ZGCDISA)

If Death checked, CONTINUE.

If Death not checked, SKIP to Question 3.

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a. Date of death (ZGCDDT)

month day year

b. Probable cause of death (ZGCDCAU)

3. Was the adverse experience: (Check all that apply)

☐ Congenital anomaly (ZGCCONA)

☐ Cancer (ZGCCAN)

☐ Life-threatening (ZGCLIFE)

☐ Due to an overdose (ZGCOVD)

IF ANY ITEM IN QUESTION B.2 OR B.3 IS CHECKED, THIS DOCUMENTS A SERIOUS ADVERSE EVENT. CONTACT THE BIOSTATISTICAL COORDINATING CENTER STAFF IMMEDIATELY AT (301) 881-9260 AND FAX THE FORM TO (301) 881-3589.

C. Event Information

1. Onset of adverse experience (ZEIONST)

☐ gradual

☐ sudden

☐ unknown

2. Was the patient on coded medication at the time of the adverse experience? (ZEICMED)

YES NO
☐ ☐

If YES, CONTINUE.

If NO, SKIP to Question 3.

a. Was the adverse experience related to coded medication? (ZEICREL)

☐ no

☐ possibly

☐ probably

☐ unknown

b. Were the coded medications interrupted or stopped? (ZEICINT)

YES NO
☐ ☐

If YES, CONTINUE.

If NO, SKIP to Question 3.

i. Which coded medication(s) was interrupted or stopped? (Check all that apply)

☐ doxazosin (ZEIINTD)

☐ finasteride (ZEIINTF)

ii. Was the adverse experience reversible when the coded medication(s) was withdrawn? (ZEICREV)

YES NO
☐ ☐

iii. Was the coded medication(s) re-started? (ZEICRST)

☐ ☐

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If YES, CONTINUE.

If NO, SKIP to Question 3.

- a) Which coded medication(s) was re-started?
(Check all that apply)

☐ doxazosin (ZEIRSTD)

☐ finasteride (ZEIRSTF)

- b) How long was the patient off coded medications?

days

(ZEIOFFN) weeks (ZEIOFFU)

months

If the patient was off doxazosin coded medications for more than 3 days, doxazosin must be re-titrated when rechallenging.

- c) Did the symptoms recur? (ZEISREC)

YES NO
☐ ☐

3. Was the patient taking any medication other than the coded medications within 14 days of the onset of the adverse experience? (ZEIMED)

☐ ☐

If YES, list below:

	Medication	Dose	Start Date	Stop Date
a.	(ZEIMEDA)	(ZEIDOSA)	(ZEISRTA)	(ZEISTPA)
b.	(ZEIMEDB)	(ZEIDOSB)	(ZEISRTB)	(ZEISTPB)
c.	(ZEIMEDC)	(ZEIDOSC)	(ZEISRTC)	(ZEISTPC)
d.	(ZEIMEDD)	(ZEIDOSD)	(ZEISRTD)	(ZEISTPD)
e.	(ZEIMEDE)	(ZEIDOSE)	(ZEISRTE)	(ZEISTPE)
f.	(ZEIMEDF)	(ZEIDOSF)	(ZEISRTF)	(ZEISTPF)
g.	(ZEIMEDG)	(ZEIDOSG)	(ZEISRTG)	(ZEISTPG)
h.	(ZEIMEDH)	(ZEIDOSH)	(ZEISRTH)	(ZEISTPH)
i.	(ZEIMEDI)	(ZEIDOSI)	(ZEISRTI)	(ZEISTPI)
j.	(ZEIMEDJ)	(ZEIDOSJ)	(ZEISRTJ)	(ZEISTPJ)

4. Describe the adverse experience in detail:

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5. Overall severity of adverse experience
(Check only one) **(ZEIOSEV)**

- ☐ 1 mild
☐ 2 moderate
☐ 3 severe

6. Duration of adverse experience
(Check only one) **(ZEIDUR)**

- ☐ 1 <1 day
☐ 2 1 day - 1 week
☐ 3 >1 week

7. Treatment administered for the adverse
experience (Check only one) **(ZEITRT)**

- ☐ 1 none
☐ 2 self-treatment / OTC drug
☐ 3 outpatient - changes in medication
☐ 4 outpatient procedure
☐ 5 inpatient hospitalization

8. Outcome (Check only one) **(ZEIOUTC)**

- ☐ 1 recovered, no residual effect
☐ 2 residual effect, no treatment
☐ 3 residual effect, being treated
☐ 4 persistent, no treatment
☐ 5 persistent, being treated
☐ 6 death

D. Conclusion

1. Additional comments:

Initials of person completing form **(FORMIN)**

first last

Form entered in computer?

☐

Signature of P.I. _____

Date _____