

Randomization Worksheet



When ready to randomize call:

1-800 253-1387 weekdays, 7am-5pm Pacific time

otherwise pager (206) 989-5212 [punch in your number (area code + phone number)]

Patient ID:

seqnum05

____-____-____

1. Today's date:

days05

____/____/____
mo day yr

Patient's name: _____

Name of person placing call: _____

Code: ____

Name of person at CTC: _____

Code: ____

qldone05

2. Is baseline Quality of Life form completed and in hand?

Yes No

If no, form not completed because
(Check all that apply)

qlref05

Patient refused

1 0

qlill05

Patient is too ill to complete it

qlilit05

Patient ~~doesn't read English~~ illiterate

qloth05

Other

qlwork05

Patient is working on form now

qlclbk05

If patient is working on form now, you will:

1 Call back for the randomization after the form is completed and in hand

2 Collect the QL form from the patient BEFORE telling the patient the assignment

3. Randomizing hospital:

_____ Code: ____

Hospital where device would be implanted

(if randomized to ICD):

_____ Code: ____

4. Initiation of AVID Therapy

If this patient is randomized to amiodarone, drug should be started within 1 day. If the patient is randomized to sotalol, either a Holter should be started within 1 day or baseline EPS scheduled within 3 days. If the patient is randomized to device, NTL implantation should be scheduled within 3 days and epicardial placement within 5 days.

Are you prepared to begin AVID therapy? strttx05

Cannot randomize unless prepared to begin.

Yes No

1 0

rtnum05

5. **Gender:** Male Female **sex00 in CIR**
1 0

6. **Race:** White Black **race05: 0=white, 1=non-white**
 Asian Native American Other

Hispanic origin: Yes No

7. **Age:** **age00 in CIR**

8. **Health Insurance:**
 Private insurance HMO VA/military
 Medicare/medicaid None

9. **QL form preference:**
 English Spanish
 Cannot participate
because: Cannot read English or Spanish
 Vision or other physical limitation
 Refuses

10. **Index arrhythmia:** Date: / /
mo day yr

arrtyp05 Type of event:
1 (A) Primary cardiac arrest due to VF, *not* associated with new MI or transient or correctable cause
2 (B) Documented sustained VT with syncope
3 (C) Documented sustained VT with near syncope, systolic BP < 80 mmHg or chest pain AND EF ≤ 0.40

inhosp05 Location of event: In-hospital Emergency Room Out-of-hospital
1 2 3

11. **Decision regarding revascularization**

Any decision regarding revascularization (includes CABG, PTCA, and atherectomy) must be made prior to randomization.

Has the decision regarding revascularization in this patient been made?

revdec05 Yes No
1 0

If "Yes", what was the decision?

revtyp05 No revascularization
4 Revascularization performed since index event
2 Revascularization planned

If revascularization (CABG, PTCA, atherectomy) performed, what was the date:

___/___/___ dycabg05
mo day yr

If revascularization (CABG, PTCA, atherectomy) planned, what date is scheduled:

___/___/___ dyrev05
mo day yr

ef00 in CIR

12. **Ejection fraction:** 0. ___ *Cannot randomize if >0.40 and arrhythmia (C) or revascularization planned.*

Date EF obtained: ___/___/___ dyef05
mo day yr

13. **Has the patient had other open heart surgery since the index event?**

othsrg05 Yes No
1 0
If yes, date of surgery ___/___/___
dysurg05 mo day yr

14. **Has signed informed consent been obtained?** infcon05 Yes No
Cannot randomize unless consent has been obtained. 1 0

15. **Are baseline forms and Registry form through item 10 complete?** blcomp05 Yes No
Cannot randomize unless forms are complete. 1 0

16. **Are any exclusions checked?** exclus05 Yes No
Cannot randomize if ANY exclusions checked. 1 0

17. Has the patient/patient's physician been advised regarding the use of aspirin, ACE inhibitors and beta-blockers? **mdadvu05** Yes No
1 0

18. Has the patient/patient's physician been informed regarding the inadvisability of changing these medications during titration of study drug? **mdadvc05** Yes No
1 0

19. Does the patient have a known contraindication to sotalol? **nosota05** Yes No
1 0
If yes, check all that apply:

- asthma05** Bronchial asthma
- brady05** Severe sinus bradycardia
- avblok05** 2-3° AV block
- hxinef05** History of clinical inefficacy
- invcon05** Investigator concern
- chfcon05** CHF
- efcon05** Low EF
- othcon05** Other specify _____
- Uncontrolled CHF **chf05**
- Hypersensitivity to sotalol **hyprsn05**
- Non-inducible or insufficient ectopy **nonind05**

20. Have you approached the patient's spouse or partner to participate in the Quality of Life study?
 Yes
 No
 Intend to soon
 Awaiting IRB approval for Spouse QL
 Other reason: _____
 Not applicable (no spouse/partner)

21. **Randomization assignment:**
iddrg00, drgtyp00 in CIR
 Device
 Drug: Amiodarone Sotalol

22. **Method of cost utilization data collection:**
 None
 "Shoebox" (accordion folder)

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