Title: Statistical Analysis Plans for clinical trials

1. Scope

1.1 This SOP applies to all clinical trials sponsored by University of Aberdeen (UoA) and/or NHS Grampian (NHSG) and applies to the statistician(s) commissioned to conduct statistical analysis and the Chief Investigator (CI).

1.2 All clinical trials must have a Statistical Analysis Plan (SAP) which is a comprehensive description of the methods and presentation of data analysis proposed.

1.4 This SOP may also be used by staff from other NHS areas, or organisations, with prior agreement.

2. Responsibilities

Chief Investigator (CI) Preparing the SAP (delegated by the Sponsor)

3. Procedure

3.1 The overall responsibility for preparing the SAP is delegated to the CI or Clinical Trials Unit (CTU) by Sponsor. This delegation of duty shall be agreed before the study begins and shall be documented in the sponsorship/site agreements.

3.2 The CI, acting on behalf of the Sponsor, may delegate the responsibility for preparing the SAP to the lead statistician but the responsibility for the final SAP remains the CIs. Any such delegation shall be documented. At least one author of the SAP shall be a professional statistician.

3.3 The SAP must be drafted, finalised and agreed before commencing the final statistical analysis of clinical trial data. Where possible statistical analysis should be complete before unblinding (revealing the treatment allocation).

3.4 Any changes to the SAP during the course of the clinical trial must be documented in the SAP and reasons for the change noted.

3.5 The SAP shall document authorship, including the statistician responsible for preparation of the

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SAP, approval from the lead statistician overseeing preparation of the SAP (if required), approval from the Chief Investigator and be version controlled.

3.6 The SAP shall state the hypotheses to be tested and/or any parameters that are to be estimated in order to meet the trial objectives stated in the trial protocol.

3.7 The SAP shall provide details of the sample size calculation reported in the trial protocol, explicitly stating all assumptions made, eg sources of data used and expert opinion sought.

3.8 The SAP shall list details of tables, figures and other data to be presented in the statistical report(s) and shall include the set of dummy data tables reflecting the contents of the final report.

3.9 The SAP shall clearly define populations within the clinical trial, for example, intention-to-treat, as randomised, sub-group analyses etc.

3.10 The SAP shall list and describe all primary and secondary outcomes, and also describe in detail any algorithms required to derive outcomes as required.

3.11The SAP shall state the frequency of interim analyses and reports.

3.12 The SAP shall include a description of the methods for analysis and presentation of the data, including, where applicable:

- Methods for point and interval estimation.
- Levels of statistical significance to be used, one-tailed or two-tailed tests to be performed, and/or clinical relevance.
- Multiple comparison methods.
- Use of baseline and covariate data.
- Methods for handling multi-centre data.
- Methods for handling missing data.
- Methods for handling withdrawals and protocol deviations.
- Identification of fixed or random effects models.
- Planned interim analysis and statistical stopping rules.
- Methods for checking critical analysis assumptions and sensitivity of assumptions.
- Rules for introduction of methods for handling missing data.
- Specification of computer systems and packages to be used for statistical analysis.

4. Abbreviations and definitions

CTU Clinical Trials Unit
SAP Statistical Analysis Plan

5. Related documentation and references

SOP-QA-3 Protocol guidance for high risk trials and CTIMPs
SOP-QA-17 Project committees
SOP-QA-18 Randomisation and blinding for controlled trials
SOP-QA-20 Data management for clinical trials
SOP-QA-21 APRs and DSURs

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