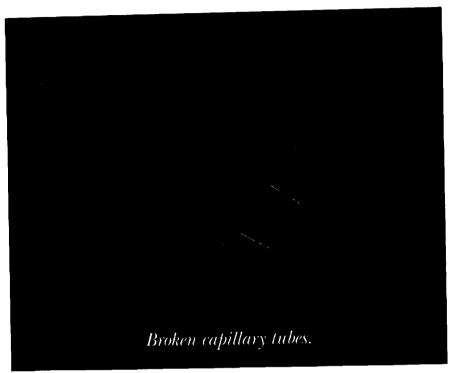
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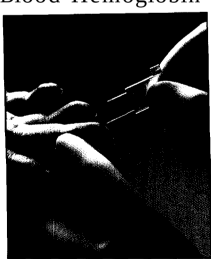
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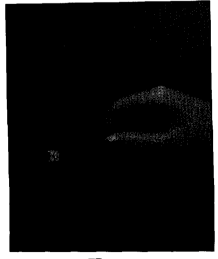
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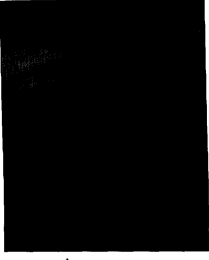


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The organization of the paper should be as follows: title page; abstract; introduction: methods; results; discussion: acknowledgments; references; tables: figures; and figure legends. The main sections and subdivisions should be indicated by side headlines flush with the left margin and two lines above the text. The Arabic numbering system should be used.

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Authors submitting manuscripts reporting the results of clinical investigations should prepare an abstract of no more than 250 words under the following headings: Objective, Design, Setting, Patients (or Participants), Interventions

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The abstract should begin with a clear statement of the precise objective or question addressed in the report. If more than one objective is addressed, the main objective should be indicated and only key secondary objectives stated. If an a priori hypothesis was tested, it should be stated.

2. Design

The basic design of the study should be described. The duration of follow-up, if any, should be stated. As many of the following terms as apply should be used.

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- b. For studies of screening and diagnostic tests: criterion standard (ie, a widely accepted standard with which a new or alternative test is being compared; this term is preferred to "gold standard"); blinded or masked comparison.
- c. For studies of prognosis: inception cohort (subjects assembled at a similar and early time in the course of the disorder and followed thereafter); cohort (subjects followed forward in time, but not necessarily from a common starting point): validation cohort or validation sample if the study involves the modeling of clinical predictions.
- d. For studies of causation: randomized control trial; cohort; case-control; survey (preferred to "cross-sectional study").
- e. For descriptions of the clinical features of medical disorders: survey; case series.
- f. For studies that include a formal economic evaluation: cost-effectiveness analysis; cost-utility analysis; cost-benefit analysis. For new analyses of existing data sets, the data set should be named and the basic study design disclosed.

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To assist readers in determining the applicability of the report to their own clinical circumstances, the study setting(s) should be described. Of particular importance is whether the setting is the general community, a primary care or referral center, private or institutional practice, ambulatory, or hospitalized care.

4. Patients or Other Participants

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2 Books

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3. Contributions to Books

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