Clinical Trial Registry

All clinical trials from India must be registered with “Clinical Trials Registry - India”. The trials conducted outside India may be registered with the respective national clinical trial registry. We have made trial registration mandatory from January 2020 for the acceptance of the study for publication.

Editorial Process

The manuscripts will be reviewed for possible publication with the understanding that they are being submitted to one journal at a time and have not been published, simultaneously submitted, or already accepted for publication elsewhere. The manuscripts are rejected by the editorial office before a formal peer-review.

The Editorial office reviews submitted manuscripts initially. Manuscripts with insufficient originality, serious scientific and technical flaws or lack of a significant message are rejected. All manuscripts received are duly acknowledged. Manuscripts are sent to two or more expert reviewers without revealing the identity of the contributors to the reviewers. Each manuscript is also assigned to a member of the editorial team, who based on the comments from the reviewers takes a final decision on the manuscript. The contributors will be informed about the reviewers’ comments and acceptance/rejection of the manuscript. The average submission to first decision time is about 3-4 weeks and about 65-70% of unsolicited manuscripts do not get published.

Articles accepted would be copy edited for grammar, punctuation, print style, and format. Page proofs will be sent to the corresponding author, which has to be returned within three days. Correction received after that period may not be included.

Authorship Criteria

Authorship credit should be based only on substantial contributions

1. Conception and design or acquisition of data or analysis and interpretation of data;
2. Drafting the article or revising it critically for important intellectual content;
3. Final approval of the version to be published.

Conditions 1, 2, and 3 must be met. Participation solely in the acquisition of funding or the collection of data does not justify authorship. General supervision of the research group is not sufficient for authorship. Each contributor should have participated sufficiently in the work to take public responsibility for appropriate portions of the content. The order of naming the contributors should be based on the relative contribution of the contributor towards the study and writing the manuscript. Once submitted the order cannot be changed without the written consent of all the contributors.

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description should be divided into the following categories, as applicable: concepts, design, the definition of
intellectual content, literature search data acquisition, data analysis, statistical analysis, manuscript
preparation, manuscript editing, and manuscript review. The author’s contributions will be printed on the
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1. **Original Articles:** Randomized controlled trials, intervention studied, studies of screening and diagnostic test, outcome studies, cost effectiveness analyses, case-control series, and surveys with high response rate. Up to 4000 words excluding about 35 reference and abstract.

2. **Review Articles:** (Including for Ethics forum, Education forum, E-Medicine, etc.) Systemic critical assessments of literature and data sources. Up to 4500 words excluding about 90 references and abstract. For review articles, include the method (literature search) in abstract as well as in the introduction section. Usually review articles are invited by the Editor-in-chief from people of eminence with vast personal experience in the field.

3. **Case Reports:** New/interesting/very rare case can be reported. Cases with clinical significance or implications will be given priority. However, mere reporting of a rare case is not encouraged and may not be considered. Up to 2000 words excluding references and abstract and up to 10 references.

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**Preparation Of The Manuscript**

**A. Title Page**

The Title page should carry

1. Types of the manuscript: Original article, Case Report
2. The title of the article, which should be concise, but informative;
3. Running title or short title, not more than 65 characters;
4. The name by which each author/contributor is known (Last name, First name, and initials of middle name) and institutional affiliation. The affiliations should be given as 1, 2, and 3 but **not** marked with symbols
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10. If the manuscript was presented as part of a meeting, the organization, place, and exact date on which it was read.
11. Registration number of clinical trials.

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The second page should carry the full title of the manuscript and an abstract (of no more than 150 words for a brief report and 250 words for original articles and other article types). The abstract should be structured for original articles and review articles. State the context (background), aims, settings and design, material and methods, statistical analysis used, results, and conclusions. Below the abstract should provide 3 to 8 keywords, arranged alphabetically. The abstract need not be structured for OR forum articles and case reports. Don’t consider references in the abstract.

**C. Introduction**

State the purpose and summarize the study or observation.

**D. Materials and Methods**

The Methods section should only include information that was available at the time the study was planned
Selection and Description of Participants: Describe your selection of the observational or experimental participants (patients or laboratory animals, including controls) clearly, including eligibility and exclusion criteria and a description of the source population. Because the relevance of such variables as age and sex to the object of research is not always clear, authors should explain their use when they are included in a study report; for example, authors should explain why only subjects of certain ages were included or why women were excluded. The guiding principle should have clarity about how and why a study was done in a particular way. When authors use variables such as race or ethnicity, they should define how they measured the variables and justify their relevance.

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Whenever possible quantify findings and present them with appropriate indicators of measurement error or uncertainty (such as confidence intervals). Report losses to observation (such as dropouts from a clinical trial). When data are summarized in the Results section, specify the statistical methods used to analyze them. Avoid non-technical uses of technical terms in statistics, such as ‘random’ (which implies a randomizing device), ‘normal’, ‘significant’, ‘correlations’, and ‘sample’. Define statistical terms, abbreviations, and most symbols. Specify the computer software used. Use upper italics (P 0.048). For all P values include the exact value and not less than 0.05 or 0.001.

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H. Discussion

Include a summary of key findings (primary outcome measures, secondary outcome measures, results as they relate to a prior hypothesis); Strengths and limitations of the study (study question, study design, data collection, analysis, and interpretation); Interpretation and implications in the context of the totality of evidence (is there a systematic review to refer to, if not could one be reasonably done here and now?, what this study adds to the available evidence, effects on patient care and health policy, possible mechanism); Controversies raised by this study; and Future research directions (for this particular research collaboration, underlying mechanisms, clinical research).

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List the first six contributors followed by et al. There should not be any gaps between the year; volume:page-page.


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3. Case report must have significant educational value including the ability to perhaps change a clinician’s traditional method of handling such a case and;
4. Case report’s interest to the reader should be significant.

**Preparation of Case Report**

Follow the standard format for the article (Abstract, Key-words, Introduction, Cases History, Discussion, and References).

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