Guide for authors

Asian Biomedicine—Guide for Authors 2018

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Aims and scope: Asian Biomedicine is an international, general medical and biomedical journal that aims to publish original peer-reviewed contributions dealing with various topics in the biomedical and health sciences from basic experimental to clinical aspects. The work and authorship must be strongly affiliated with a country in Asia, or with specific importance and relevance to the Asian region. The Journal will publish reviews, original experimental studies, observational studies, technical and clinical (case) reports, practice guidelines, historical perspectives of Asian biomedicine, clinico-pathological conferences, and commentaries.

Innovative investigations including basic research; clinical trials; studies of diagnostic accuracy; studies relating to behavioral, therapeutic, or epidemiological aspects of medicine; public health, clinical guidelines, standards of health care, and indigenous diseases are welcomed. We also welcome manuscripts about the history of Asian medicine, and controversies. We invite guest editorials on occasion.

Audience: Asian biomedicine is intended for a broad and international audience, primarily those in the health professions including researchers, physician practitioners, basic medical scientists, dentists, educators, administrators, those in the assistive professions, such as nurses, and the many types of allied health professionals in research, and health care delivery systems, including those in training.

The Editors of Asian Biomedicine require authors to be in compliance with the recommendations of the International Committee of Medical Journal Editors (ICMJE) for the conduct and reporting of scholarly work in medical journals. The current recommendations (December 2017 at printing) are available at http://www.icmje.org. The Editors of Asian Biomedicine adhere to principles of research integrity and aim to avoid any type of scientific misconduct, such as fabrication, falsification, plagiarism, redundant publication, and authorship problems. All submitted manuscripts are checked for potential plagiarism of all types, including patchwork plagiarism, using specialist services including those from iThenticate, Crossref, and ProQuest, and are reviewed by an editor responsible for publication integrity. In resolving any potential scientific misconduct, Asian Biomedicine follows international standards, guidelines, and flowcharts provided by the Committee on Publication Ethics (COPE) (available at: http://publicationethics.org/resources), the Council for International Organizations of Medical Sciences (CIOMS), the World Association of Medical Editors (WAME) (http://www.wame.org/about/recommendations-on-publication-ethics-policy), and the Council of Science Editors (http://www.councilscienceeditors.org/resource-library/editorial-policies/). Asian Biomedicine gives equal consideration to every carefully performed study investigating an important question within the scope of the Journal, whether the results are negative or positive.

This Guide for Authors is revised periodically by the Editors as needed, and includes our ethical and policy guidelines. Authors should consult a recent issue of the Journal or visit www.asianbiomed.org for the latest version of these guidelines. Any manuscript not prepared according to these guidelines may be returned to the author(s) for revision without review.

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Index

1 Manuscript submission
   1.1 Submission
   1.2 Important information
   1.3 Supporting documents
2 Disclosure of conflicts of interest
3 Human and animal rights, and ethical considerations
4 Reporting clinical trials
5 Informed consent for publication
6 Previous publication or duplicate submission
7 Article categories
   7.1 Reviews
   7.2 Minireviews
   7.3 Practice guidelines
   7.4 Original articles
   7.5 History of Asian biomedicine
   7.6 Clinical reports
   7.7 Clinical vignettes
   7.8 Technical reports
   7.9 Brief communications
   7.10 Controversies in biomedicine
   7.11 Commentary
   7.12 New developments
   7.13 Clinicopathological conferences
   7.14 Letters to the Editor
8 Manuscript preparation
   8.1 Title page
   8.2 Article title
   8.3 Abstract and Keywords
   8.4 Main text
      8.4.1 Abbreviations
      8.4.2 Jargon, slang, colloquialisms, idioms, homonyms, euphemisms, and clichés
   8.4.3 Numbers
   8.4.4 Units
   8.4.5 Names of drugs, devices, and other products
   8.4.6 Gene nomenclature
   8.4.7 Genomic and related data
   8.4.8 Plant (or other) extracts
   8.4.9 Cell lines
   8.4.10 Blots and gels
   8.4.11 Antibodies
   8.4.12 Tumor cell lines and anticancer activity
   8.4.13 Statistical requirements
   8.4.14 Qualitative research
   8.4.15 Personal communications and unpublished data
   8.5 Authors’ contributions
   8.6 Conflict of interest statement
   8.7 Acknowledgments
   8.8 References
      8.8.1 In the main text, Tables, Figure legends
      8.8.2 In the references list
   8.9 Tables
   8.10 Figures
      8.10.1 General guidelines
      8.10.2 Formats
   8.11 Manuscript checklist (before submission)
9 The editorial and peer review process
10 Preparation for publication
11 Publication charges and reprints
12 Copyright
13 Permissions
1 Manuscript submission

1.1 Submission

All submissions must be made online via the Editorial Manager submission system. (https://www.editorialmanager.com/abm/Default.aspx). This system is generally intended to be intuitive, but an Author Tutorial is found by clicking on the “here” hyperlink for “Additional help with your submission,” please click here for the Author Tutorial” on the Editorial Manager Author Main Menu page after logging in as an author. Alternatively, the Tutorial for Authors is under the “Instructions for Authors” hyperlink in the white on black menu at the top of the page. An exception to the Tutorial for Authors is that we do not charge submission or other fees and so the section on “Fees and Payments” does not apply. The electronic manuscript (meaning all submission items, including all text, Tables, Figure artwork, conflict of interest disclosures, and any other required documents) should be uploaded through the Editorial Manager system found under the Submit button on the Journal website (http://www.asianbiomed.org). Please do not submit manuscripts by e-mail, post, or fax, or hand deliver your manuscripts to the Editorial Office.

If you do not receive an acknowledgment from the Journal within 10 days or assistance is required, please contact the Editorial office. With any communication with Asian Biomedicine please include the assigned manuscript identification number once provided.

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1.2 Important information

- Manuscripts should be uploaded in Microsoft Word document (not PDF) format and prepared in the simplest form possible. The text should be left aligned, not justified. Please use a typeface or font with high legibility, such as Times New Roman, 12 point, on A4 pages with 2.54 cm (1 inch) margins. After acceptance, we will format the manuscript according to the Journal’s style. We may also modify formatting to facilitate peer review.
- Articles should meet the following basic criteria: the information is important, the writing clear and concise, the study methods appropriate, the data valid, and the conclusions reasonable (not overstated) and supported by the data.
- Number the pages consecutively. You may use automatic page numbering, but do NOT use other kinds of automatic formatting, such as for footnotes, headers and footers, or reference lists. Reference lists prepared using bibliographic software such as EndNote should have the field codes removed (select all (cntl+a) and then use ctrl+shift+f9 on the keyboard).
- Put the main text, references, Table headings, and Figure legends in one file.
- Tables (as editable text in an MS Word document) and Figures should be submitted as separate files. For peer review, most formats and styles are acceptable. However, they must be prepared according to our production standards on revision and before acceptance of the manuscript for publication, and such revision before submission may facilitate peer review. To facilitate the editorial process and uniquely distinguish your files from the many others received, please name files according to the figure number and identify with the name of the corresponding author and manuscript reference number if assigned, e.g., “00035_AuthorName_Fig1.tif”, “00035_AuthorName_Fig2.jpg.” Please see section 8 of this guide.
- Manuscripts must be written in acceptable U.S. American English. Stedman’s Medical Dictionary (27th edition or later), the American Medical Association Manual of Style (9th edition or later), and Merriam-Webster’s Collegiate (10th edition or later) or Unabridged Dictionary should be used as standard references. Manuscripts that are accepted for publication will be checked by our copyeditors and edited according to the Journal’s house style. If English is not your native language, or substantial English language and style editing is required, we normally ask that you have your manuscript checked by a native English speaker, preferably with knowledge of your field. Clear and concise language, and appropriate grammar and word choice will help the Editors and reviewers concentrate on the scientific content of your manuscript so that it can be judged fairly on its merits and thus facilitate the
peer review process. If the use of language is poor, the manuscript is unlikely to attract a dispassionate or unbiased peer review. If you do not know a native English speaker with knowledge of your field, please consider using a professional English language editing service to help prepare your manuscript before submission, but note that the use of such services is no guarantee of acceptance. Editors hold the right to make any changes to the language and style of the original manuscript to adhere to the uniform standards of Asian Biomedicine, and authors may review these changes in proof.

1. A professional e-mail address (from an academic institution with an appropriate second-level domain such as .ac or .edu or institutional) is usually required for the corresponding author (publication of Gmail, Yahoo, Hotmail, or other free e-mail addresses alone will not usually be acceptable) and is preferred for other authors. All Editorial correspondence concerning receipt, status, review, revision, and publication of a manuscript will be sent to only one person who has been designated as the corresponding author during submission and evaluation. The corresponding author is responsible for communicating the manuscript status to all coauthors and for obtaining the coauthors’ assent for any substantial changes of content or interpretation made during revision. We encourage the use of an ORCID, especially for corresponding authors. Please see the following URL for further details, https://orcid.org/. An ORCID provides a persistent digital identifier for contributors.

1.3 Supporting documents

The following documents or statements should generally be uploaded if relevant to your submission (refer also to the Manuscript checklist that follows these guidelines). Items (2), (3), (9), and (10), are mandatory. Items (1), (4), (5), (6), (7), and (8) are required only if they apply to your manuscript. Except for the Open Access License, these documents are not required for editorial consideration of the manuscript. However, they may be required for peer review or further processing, and where mandatory will be required for acceptance for publication.

1. A cover letter is no longer required because most of the required information will be uploaded or provided during the online submission process. At the Comments stage of the submission process, please enter any such additional comments you would like to send to the publication office. These comments will not appear directly to reviewers in your submission. However, it may facilitate the editorial process if you include a short description of the research or topic of the manuscript (2–3 sentences) and briefly any other information regarding the manuscript that the Editors may find useful, such as reasons why authors presume their work may be of interest to the readership of Asian Biomedicine, and a list of manuscripts that have been published, submitted, or are in press that are similar to the submission to Asian Biomedicine. Please include in your submission, copies of those similar manuscripts that might be regarded as redundant publications so that the Editors can be assured there is no redundancy.

2. Authorship statement. Each author’s contribution to the manuscript should be noted as described in the Authorship Statement form available at the home (welcome) page of Editorial Manager for Asian Biomedicine. These contributions should also be included in an author contributions addendum at the end of the manuscript. Unless there are exceptional circumstances, the signatures of ALL coauthors must be included on the authorship statement or the manuscript cannot be accepted for publication. Please note that copied and pasted “graphics” of signatures or “e-signatures” are NOT acceptable because of the possibility of fraud. In cases where obtaining a signature from each author would unduly delay publication, the corresponding author’s signature is sufficient provided that the corresponding author understands that he or she signs on behalf of the other authors who have not signed the form and has their consent to do so. If it is not possible to obtain the signature of a particular author (owing to death, loss of contact, or other reasons), the corresponding author should sign in their place and explain the circumstances for each instance in a signed document.

Changes to authorship: the author list of any submission should be decided upon and fixed BEFORE submission. Other than only in exceptional circumstances, the Journal does not allow addition or removal of author names after submission. A satisfactory explanation for any proposed changes in authorship will be required and ALL authors will be required to supply a new signed consent form that reflects the changes. We will also require a signed consent form from any person whose name has been removed indicating that they agree to the removal of their name from the author list. Because of the complexity of such situations, we strongly advise authors to fix the author list before submission and not to attempt to make changes later.

For authorship please follow the 4 criteria recommended by the ICMJE: 1. “Substantial contributions to the conception or design of the work; or the acquisition, analysis, or interpretation of data for the work”; AND 2. “Drafting the work or revising it critically for important intellectual content”; AND 3. “Final approval of the version to be published”; AND 4. “Agreement to be
accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.” The 4th criterion is covered in the statement on the signature page. The other 3 criteria are covered in the authorship statement. An author is considered to be someone who has made substantive intellectual contributions to a published study and must take responsibility for at least one component of the work. They should be able to identify who is responsible for each other component, and should ideally be confident of their coauthors’ ability and integrity. “Gift” authorship is not acceptable.

• We do note that the ICMJE authorship criteria stated on the form do not exactly match all of the CRedit Taxonomy for Contributor Roles that should be entered in the online Editorial Manager system during submission. We suggest that the roles that most closely match the criteria are as follows. **Criterion 1.** Conception and design: Conceptualization, Funding acquisition, Methodology, Project administration, Supervision. 1. Acquisition of data: Data curation, Investigation, Software, Resources, Visualization. 1. Analysis and interpretation of data: Formal analysis, Validation. **Criterion 2.** Drafting the manuscript: Writing – original draft. 2. Critically revising the manuscript: Writing – review & editing. **Criteria 3 & 4.** Check the box marked Other and enter the role “Approved the final version and take responsibility for statements made in the published article.”

• The corresponding author is responsible for obtaining the approval of other authors of the original submission and any subsequent revision, and for all correspondence accompanying the manuscript. In the published article, at least one, or optionally more than one author must be designated with an asterisk as the author(s) to whom reader correspondence regarding the published article may be addressed. This does not need to be the same as the corresponding author, but ideally should be somebody whose contact details are not likely to change in the near future and must except in unusual circumstances have a professional or institutional email address. We appreciate those in supervisory or administrative roles may be more senior authors who may not have the time to communicate with journal editors, and other authors. They may also not have sufficient time, experience or patience for the online submission processes.

• Present addresses or affiliation for individual authors that differ from the address(es) at which the work was done should be given in a Present Address or Affiliation note in the Acknowledgments section and this should be highlighted in the Comments section during the submission process. Please see our Guide for Authors section 8.1.

• All individuals who are listed as authors must have read and approved the final version of the manuscript. This requirement is the responsibility of the corresponding author. If it is not possible for a particular author to read and approve the final version of the manuscript (owing to death, loss of contact, or other reasons), the corresponding author explain the circumstances for each instance in a signed document.

• All people who have made substantial contributions to the work, but do not meet the 4 criteria for authorship and thus do not fulfill the requirements to be listed as authors, should be listed in the Acknowledgments section. Without other contributions, the acquisition of funding, general supervision of a research group, or general administrative help, provision of materials, collection of data, technical help, writing assistance, or English language editing, alone do not usually justify authorship. The contributions made by people listed in the acknowledgments should be specified (e.g., “served as scientific advisors”, “critically reviewed the study proposal”, “collected data”, “provided and cared for study patients”, “assisted in writing and technical editing of the manuscript”). Because acknowledgment may imply endorsement of an article by the acknowledged individuals, all persons named in the Acknowledgments section of the manuscript must give their permission in writing to be named. We do not normally need to see this written approval, but it should be stored safely in case it is ever required. The involvement of scientific (medical) writers or anyone else who assisted with the preparation of the manuscript content should be acknowledged, along with their source of funding, as described in the commentary by Jacobs and Wager. European Medical Writers Association (EMWA) guidelines on the role of medical writers in developing peer-reviewed publications. Current Medical Research and Opinion. 2005; 21(2): 317–321, found at http://www.emwa.org/documents/about_us/EMWAGuidelines.pdf. The role of medical writers should be explicitly acknowledged in the Acknowledgments or Authors’ contributions sections as appropriate so that such contributions are acknowledged and transparent. Acknowledgments should be brief, and should not include thanks to anonymous referees and editors, or effusive comments. Funding including grant numbers can be included in this section. If no specific funding was received please write “We did not receive any specific grant for this research from any funding agency in the public, commercial, or not-for-profit sectors.”

• During the online submission process the corresponding author will be required to affirm that anyone who participated substantially in the study has not been omitted from
3. Statement of originality. During the online submission process, the corresponding author must declare that the material contained in the manuscript submitted, including data, text, Figures, and Tables are original and have not been published previously in the same or similar form (except as an abstract of a meeting presentation or preliminary report, e.g., on a clinical trials registry, which should be noted in the Acknowledgments section), and must not be under consideration for publication elsewhere. No separate document is required.

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5. Ethics statement. For manuscripts covering the use of human participants, human data, animals (vertebrates or any regulated invertebrates), or (in most cases) biological samples, such as clinical isolates, the letter of approval from the relevant ethics review committee should be uploaded. See Section 3.

During the online submission process the corresponding author must provide a formal statement of ethical conduct and state whether the reported research was approved by an institutional/national ethics committee or review board. If so, authors will need to state the name of the approving committee or review board and approval reference number where appropriate. If not, reasons for the exception must be provided.

A statement detailing this, including the name of the approving institutional review board or ethics committee and the reference number, must appear in all manuscripts reporting such research. If a study has been granted an exemption from requiring ethics approval, this should be detailed in the manuscript (including the name of the ethics committee that granted the exemption). If a study has not been granted ethics committee approval before it was started, retrospective approval cannot usually be obtained and it will not be possible to consider the manuscript for peer review.


7. Statement of documented informed consent. The Editors may require proof of appropriate documented informed consent from each research participant in studies involving humans. Do not send original documents or copies to the journal, but keep them safely archived, for access should they ever be required. During the online submission process the corresponding author is required to provide a formal statement that appropriate documented informed consent was obtained from each research.

the authorship list and that all persons listed as authors qualify for authorship. Furthermore, that all persons who have made substantial contribution to the work, but do not meet the authorship criteria are listed in the Acknowledgments section (technical help, writing assistance, general support, financial and material support) and that all persons named in the Acknowledgment section of the manuscript have given their permission in writing to be named (do not send this, but keep it safely archived in case required).

- Group authorship: when a large multiauthor group has conducted the work, the group ideally should decide who will be an author before the work is started and confirm who is an author before submitting the manuscript for publication. All members of the group named as authors should meet all 4 criteria for authorship, including approval of the final manuscript, and they should be able to take public responsibility for the work and should have full confidence in the accuracy and integrity of the work of other group authors. They will also be expected as individuals to complete ICMJE forms for disclosure of any potential conflict-of-interest. Some large multiauthor groups designate authorship by a group name, with or without the names of individuals. When submitting a manuscript authored by a group, the corresponding author should specify the group name if one exists, and clearly identify the group members who can take credit and responsibility for the work as authors. The byline of the article identifies who is directly responsible for the manuscript, and PubMed lists as authors whichever names appear on the byline. If the byline includes a group name, PubMed will list the names of individual group members who are authors or who are collaborators, sometimes called nonauthor contributors, if there is a note associated with the byline clearly stating that the individual names are elsewhere in the paper and whether those names are authors or collaborators. If you would like the names of the individual members of a collaboration Group to be searchable through their individual PubMed records, please ensure that the title of the collaboration Group is included on the title page and include collaborating author names as the last paragraph of the Acknowledgments section. Please add authors in the format: first name, middle initial(s) (optional), and last name. You can add institution or country information for each author if you wish, but this should be consistent across all authors.
2 Disclosure of conflicts of interest

A conflict of interest occurs when an individual’s objectivity is potentially compromised by a desire for financial gain, prominence, professional advancement, or a successful outcome. The Editors of Asian Biomedicine endeavor to ensure that what they publish is as balanced, objective, and evidence-based as possible. The potential for a conflict of interest (also known as a dual commitment, competing interest, or competing loyalty) can exist whether or not an individual believes the relationship affects their judgement. Because it is difficult to distinguish between an actual conflict of interest and a perceived conflict of interest, Asian Biomedicine requires authors to disclose all and any potential conflicts of interest.

Conflicts of interest may be financial or nonfinancial. Financial conflicts are the most easily identifiable, and include financial relationships such as payments of honoraria such as for attendance at meetings, educational grants, participation in speakers’ bureaus, membership, employment, consultancies, stock ownership, or other equity interest, expert testimony, or patent-licensing arrangements. Nonfinancial conflicts are more nebulous, and include personal or professional relationships, affiliations (other than those affiliations listed in the title page of the manuscript), academic competition, intellectual passion, ideology, or knowledge or beliefs that might affect objectivity. The conflicts can vary from those with negligible potential to those with great potential to influence objectivity and judgement. However, not all relationships represent a true conflict of interest.

Authors must disclose any financial interests or other conflicts that might, or might be perceived, to influence the conclusions or outcome of studies reported. Financial support from a commercial source, or support for travel to conferences by industry, must be stated. If a study is funded by an agency with a proprietary or financial interest in the outcome of the study, the corresponding author must include the following statement in the manuscript upon submission: “I had full access to all of the data in this study and I take complete responsibility for the integrity of the data and the accuracy of the data analysis.” Authors from pharmaceutical companies, or other commercial organizations that sponsor clinical trials, should declare this in the Conflicts of Interest statement as a competing interest on submission. To ensure

The International Committee of Medical Journal Editors (ICMJE) has developed a Form for Disclosure of Potential Conflicts of Interest to facilitate and standardize authors’ disclosures. The form is available for download at the home (welcome) page of Editorial Manager for Asian Biomedicine or from http://icmje.org. Please follow the instructions to complete the interactive e-form carefully, including the use of Adobe software such as Reader, because the form will not work with browsers and other software. Please use the interactive form to generate a disclosure statement for each author using the button in section 6. The corresponding author should collect the conflict of interest disclosure forms completed according to the ICMJE instructions from all authors. In author collaborations where formal agreements for representation allow it, such as written agreement from other authors and where the rules of their institutions allow; it is sufficient for the corresponding author to complete a form on behalf of other authors, but we must have a completed form for each author. A Conflicts of interest statement will be generated for the accepted version of the manuscript identifying any potential conflict of interest disclosures or not, but authors should also include conflicts of interest disclosures in the submitted manuscript. For accepted manuscripts, the corresponding author will be asked to confirm that each of the author’s disclosures or declarations of no potential conflicts of interest are accurate, up-to-date, and consistent with those reported in the manuscript.

The following summarizes some key considerations that must be satisfied for publication in Asian Biomedicine.

For investigations involving human participants or data, or human material samples, or animal studies or samples, appropriate institutional review board or ethics committee approval is required, and such approval must be stated in the Methods section of the manuscript. Research involving human subjects, identifiable human samples (such as urine, blood, serum, or tissue), and personal and health record data must be subjected to a review by a formally constituted institutional ethical review board. Studies must in any case be in accordance with the principles outlined in the contemporary revision of the Declaration of Helsinki of 1964 (World Medical Association (WMA) incorporating the most recent (October 2013) and earlier amendments. The Declaration of Helsinki: ethical principles for medical research involving human subjects, is available at: http://jama.jamanetwork.com/article.aspx?articleid=1760318 and http://www.wma.net/en/30publications/10policies/b3/). Any doubtful aspects of compliance with the principles of the contemporary revision of the Declaration of Helsinki must be explained and explicitly approved by the investigators’ institutional review board. For those investigators who do not have formal institutional ethics review committees, the contemporary revision of the Declaration of Helsinki and recommendations of the ICMJE must be followed.

For investigations in humans, indicate in the manuscript whether the procedures followed were in accordance with the contemporary (current) revision of the Declaration of Helsinki, state explicitly in the Methods section of the manuscript that informed consent was obtained from all participating adults, or from the parents or legally authorized guardians or representatives of children (minors or less than 18 years old) or incapacitated adults, together with the way the informed consent was obtained (explicitly, verbal or written) and documented. Evidence of informed consent by participants and respect for the privacy of the study subjects is important. The authors must be able to produce documented evidence of informed consent and respect for privacy of the study subjects if required. Because patient confidentiality is better guarded, we request that authors archive the documented informed consent and provide Asian Biomedicine with a written statement that attests that they have received and archived documented patient consent. An archive period of 5 years after publication is considered standard, and perpetual archiving is desirable. Please note that manuscripts regarding humans that may be identified must be accompanied by a documented (preferably signed) statement of informed consent to publish (in print and online) the descriptions, photographs, or pedigrees from each subject who may be able to be identified (Please see section 5). Please note

### 3 Human and animal rights: ethical approval of studies and informed consent to be included in studies

The ethical guidelines for publication in Asian Biomedicine are to ensure that the Asian Biomedicine review process selects manuscripts for publication that satisfy high quality standards while preventing exploitation of humans and animals for research. The Asian Biomedicine process is thorough, objective, fair, and conforms to international ethical standards specified in the introduction and elsewhere in these instructions.
that this is required for all case reports and series, whether or not the patient may be identifiable.

If minors are old enough to understand the proposed research, its potential risks and possible benefits, and the role expected of them as participants, then it is also desirable to obtain their consent. Assent means to express willingness to participate in research by individuals who are by legal definition too young to give informed consent. However, assent by itself is not sufficient. Even if assent is given, informed consent must still be obtained from the participant’s parent or legal guardian. Legal definitions of who constitutes a “minor” dictate whether or not a person can consent to participate in a protocol. The State of Maryland, USA, considers a person 18 years and older to be an adult and therefore one who can provide consent without parental permission. In the absence of a local older age requirement, we elect to follow Maryland law. Assent may be obtained by talking with the child and supporting that talk with a written assent document (e.g. form NIH-2514-2) appropriate to the child’s age and level of comprehension. The parents or legal guardian of a child participant should be fully informed and given a protocol consent document to read and sign before their child participates in the protocol. If the child cannot read and oral assent is obtained from the child, this should be documented on the consent document. A prewritten statement on the signature page of the consent document should read, “The information in the above consent was described to my child and my child agrees to participate in the study.” The parent’s or guardian’s signature indicating consent is in addition to the one confirming the statement of assent.

Clinical isolates and biobank samples. The consent of participants is sometimes, but not always, required before biobank samples and clinical isolates can be used in research. The use of samples must at least conform to the U.S. Office for Human Research Protections (OHRP; Rockville, MD, USA) in the U.S. Department of Health and Human Services (DHHS) regulations for the protection of human subjects (45 CFR part 46) and the additional ethical principles in the WMA Declaration of Taipei on Ethical Considerations regarding Health Databases and Biobanks, adopted by the 53rd WMA General Assembly, Washington, DC, USA, October 2002 and revised by the 67th WMA General Assembly, Taipei, Taiwan, October 2016. These regulations and principles require consent to use samples and/or data that are identifiable, and so exclude those samples that have been coded. The Declaration of Helsinki, states that “Medical research involving human subjects includes research on identifiable human material or identifiable data” and must be approved by an institutional ethics review board (IRB) or a research ethics committee (WMA, 2013). Any research using nonidentifiable samples is allowed without the need to obtain informed consent and approval of the protocol from an IRB or a research ethics committee. Nonidentifiable or anonymized biological material is that stored alongside associated information, for example, the type of tumor, medical treatment, donor’s age and so forth, but all information that would allow identification of the research participant or patient is stripped, either irreversibly (unlinked anonymized) or reversibly (linked anonymized). In the case of linked anonymized samples, identification is still possible by a code, to which researchers or other users of the material—as part of the definition of the term reversibly linked anonymized—do not have access. Coded samples have the same characteristics as linked (reversibly) anonymized samples, the only difference being that researchers and users have access to the code. Finally, samples are considered to be identified if the information that allows identification—name, address and so on—is associated directly with the tissue, such as when the patient’s name tag is attached to the sample. This is, for example, how pathology departments usually store clinical samples. The matters are complex and further guidance can be obtained from Elger and Caplan. Consent and anonymization in research involving biobanks. European Molecular Biology Organization (EMBO) reports. 2006; 7:661–6.

Clinical isolates, any bacterial or fungal subcultures, and virus isolates, made from clinical specimens, are not part of the specimen themselves, and are considered not to belong to the patient from whom they were obtained. According to Subpart A of the DHHS regulations, also called the Common Rule, if the study did not involve an investigator interacting or intervening with living individuals for research purposes to obtain the isolates, and the investigator does not obtain identifiable private information about those living individuals, then it is our view that the investigator is not conducting research on human subjects under the U.S. Department of Health and Human Services regulations for the protection of human subjects 45 CFR 46.102(f), and thus the study would not require IRB review (see http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html#46.102).

Asian Biomedicine will not accept manuscripts whose data derives from transplants involving organs obtained from executed prisoners. Asian Biomedicine requires all procedures and studies involving human participants to have been conducted according to the ethical guidelines outlined by The Transplantation Society at http://www.tts.org and have involved no use of organs or other material from executed prisoners, or other unethical practices in obtaining donor organs. It is a fundamental principle of The Transplantation Society that organs and tissues are given freely and without coercion. As a consequence, presentations of studies involving
patient data or samples from recipients of organs or tissues from executed prisoners cannot be accepted. A prerequisite for accepting a manuscript is that the authors’ work complies with the current Declarations of Helsinki and Istanbul (www.declarationofistanbul.org).

In addition, scientific studies and clinical activities should be performed in keeping with the ethical principles delineated in the following policy documents:


For work involving animals including vertebrates or any regulated invertebrates, full compliance with local, national, and international regulations is necessary, and the protocols must be reviewed in advance by the relevant institutional ethical review board for the care and use of laboratory animals. Procedures involving any animal are to be undertaken only with the goal of advancing scientific knowledge and with the explicit approval of an Institutional Animal Care and Use Committee (IACUC) before they begin. When reporting experiments involving animals, indicate in the manuscript whether the procedures followed were in accordance with the relevant regulations for the care and use of animals. The guidelines for their care and use that were followed, and any relevant licences, should be stated in the Methods section of the manuscript. At least U.S. guidelines must be followed. All animal experiments must conform to the revised Institute of Laboratory Animal Resources, Commission on Life Sciences, National Research Council “Guide for the Care and Use of Laboratory Animals” Washington, D.C.: National Academy Press; 1996. These documents can be obtained at: http://grants.nih.gov/grants/olaw/olaw.htm and the same should be stated in the Methods section of the manuscript. The European Commission Directive 2010/63/EU revising Directive 86/609/EEC for animal experiments (available at http://ec.europa.eu/environment/chemicals/lab_animals/legislation_en.htm), the revised Animals (Scientific Procedures) Act (ASPA) 1986 in the UK (https://www.gov.uk/government/publications/operation-of-aspa), or The Animals for Scientific Purposes Act, BE 2558 (AD 2015), and the manuscript should require the death of an animal must employ the most humane euthanasia method that is consistent with the goal of the study and the recommendations of the Report of the American Veterinary Medicine Association Panel on Euthanasia. J Am Vet Med Assoc. 2001; 218:669–96. The Editors reserve the right to reject any manuscript containing studies that does not or appears not to conform to these recommendations for the use of animals in research or for which procedure approval documentation cannot be provided. Please provide a manuscript prepared according to the ARRIVE guidelines checklist available at https://www.nc3rs.org.uk/arrive-guidelines.

Any field studies and other nonexperimental research on animals must comply with institutional, national, or international guidelines, and where available should have been approved by an appropriate ethics committee, which is a requirement in Thailand since the Animals for Scientific Purposes Act, BE 2558 (AD 2015), and the manuscript should include a statement specifying the appropriate permissions and/or licences. We require that authors comply with the Convention on Biological Diversity (http://www.cbd.int/convention/) and the Convention on the Trade in Endangered Species of Wild Fauna and Flora (http://www.cites.org/).

4 Reporting clinical trials

In strict accordance with ICMJE recommendations and the Declaration of Helsinki, Asian Biomedicine requires as a condition of consideration for publication that all clinical trials be registered with a database that is readily available to the public. Public trials registries include http://www.clinicaltrials.gov/ and other registries acceptable to the ICMJE (for a list see http://www.icmje.org/about-icmje/faqs/clinical-trials-registration/), such as the Thai Clinical Trials Registry (TCTR) (http://www.clinical.trials.in.th), and any other of the primary registries in the WHO Registry Network that
participate in the WHO International Clinical Trials Registry Platform (for a list see http://www.who.int/ictrp/network/primary/en/). Please include the Clinical Trial Registration number and registry name in the Abstract and Methods section. Include in the Methods section a statement that the study followed the current principles of the Declaration of Helsinki, that subjects gave their documented, informed consent; and that the study was approved by an institutional ethics review board. Please note that the date of registration must precede or be concurrent with the date of selection of the first participant. Trials without a verifiable registration number will not be considered for peer review or publication. Please see http://www.icmje.org/recommendations/browse/publishing-and-editing-issues/clinical-trial-registration.html for further detail.

The purpose of clinical trial registration is to prevent selective publication and selective reporting of research outcomes, to prevent unnecessary duplication of research effort, to help patients and the public know what trials are planned or ongoing into which they might want to enroll, and to help give ethics review boards considering approval of new studies a view of similar work and data relevant to the research they are considering. Retrospective registration, for example at the time of manuscript submission, meets none of these purposes.

All randomized controlled trials submitted for publication should upload a completed Consolidated Standards of Reporting Trials (CONSORT) flow chart (please go to http://www.consort-statement.org for more information), which provides a standard way for researchers to report trials, allowing reviewers and readers to evaluate their validity more appropriately. Other sources of reporting guidelines are the EQUATOR Network (www.equator-network.org/home/) and the NLM’s Research Reporting Guidelines and Initiatives (www.nlm.nih.gov/services/research_report_guide.html).

The ICMJE use as the definition of a clinical trial: “any research project that prospectively assigns people or a group of persons to an intervention, with or without concurrent comparison or control groups, to study the cause-and-effect relationship between a health-related intervention and a health outcome. Medical interventions include drugs, surgical procedures, devices, behavioral treatments, process-of-care changes, and the like.” This definition includes phase I to IV trials. The ICMJE define health-related interventions as “any intervention used to modify a biomedical or health-related outcome” and health-related outcomes as “any biomedical or health-related measures obtained in patients or participants.” If you are unsure whether your trial needs registering, see the ICMJE FAQs for further information. Purely observational studies (those in which the assignment of the medical intervention is not at the discretion of the investigator) do not require registration, but should be reported following the STROBE statement and its extension as appropriate, for observational studies in conjunction with the explanatory article (see von Elm, Altman, Pocock, Gøtzsche, Vandenbroucke. STROBE Initiative. The Strengthening the Reporting of Observational Studies in Epidemilogy (STROBE) statement: guidelines for reporting observational studies. Bull World Health Organ. 2007; 85:867–72 or http://strobe-statement.org).

5 Informed consent for publication of identifying details of patients in descriptions, photographs and pedigrees

“Patients have a right to privacy that should not be violated without informed consent” (ICMJE recommendations, December 2017 and earlier versions). It is necessary for authors to ensure that that the anonymity of patients is carefully protected. The names, initials, hospital or national identity numbers, or dates of birth of patients and research subjects should not be used. Other personal or identifying information should not be used unless it is essential to the clinical message or scientific purpose of the article and the patient (or parent, or legally authorized guardian or representative for minors or incapacitated adults) gives written informed consent for publication. A signed statement of informed consent to publish (in print and online) patient descriptions, photographs, or pedigrees should be obtained from all persons (or parents, or legally authorized guardians or representatives) who can be identified in such written descriptions, photographs, or pedigrees (including by the patients themselves). Informed consent for this purpose requires that such persons should be shown the manuscript before its submission or have waived the opportunity to do so in writing.

Even where consent has been given, identifying details should be omitted if they are not essential. At the discretion of the Editors, omitting data or making data less specific to deidentify patients may acceptable if alterations do not distort scientific meaning, but changing any such data is not acceptable. Previously used measures to attempt to conceal the identity of an individual in a photograph, such as placing black bars over the person’s eyes or blurring the face of the individual concerned, are not effective. Individuals can be identified in photographs that show minimal body parts, usually from identifying features (e.g. a tattoo, jewelry, clothing, fancy nail polish, scars, a nevus, or a mole). To avoid identifiability in such cases, photographs should be cropped. However, complete anonymity is difficult to achieve, and informed consent should be obtained if there is any doubt whatsoever. Please...
note that this is required for all clinical (case) reports and series, whether or not the patient may be identifiable. On submission please optionally use the Patient privacy and confidentiality form available at the home (welcome) page of Editorial Manager for Asian Biomedicine. Please note that a completed form is mandatory and must be supplied before peer review. In the Methods section, please state “(each) patient was given an opportunity to review the manuscript and provided documented consent for its publication.”

Nonidentifiable images

At the discretion of the Editors, formal consent is not usually required for the use of entirely anonymized images from which the individual cannot be identified, e.g., x-ray images, ultrasound images, intraoperative images, pathology slides, or laparoscopic images, provided that these do not contain any identifying marks and are not accompanied by text that might identify the individual concerned. Nevertheless, consent on our Patient privacy and confidentiality form available at the home (welcome) page of Editorial Manager for Asian Biomedicine is required for all clinical (case) reports and series regardless of whether the patients can be identified or not.

6 Previous publication or duplicate submission

Submitted manuscripts are considered with the understanding that they have not been published previously in print or electronic format (except in abstract or poster form) and are not under consideration in totality or in part by another publication or electronic medium. Such a statement of originality will be required as part of the corresponding author questionnaire to be completed during online submission of work through the Editorial Manager system. Published abstracts or presentations at conferences should be noted in the Acknowledgments section of the manuscript. Asian Biomedicine follows the recommendations of the ICMJE with regard to overlapping publications and acceptable secondary publication, including translations of work published in a language other than English or multiple publications based on the same database.

Authors should be aware that replication of text from their own previous publications is text recycling (also referred to as self-plagiarism), and is mostly considered unacceptable. Where similarity of text with authors’ own previous publications is necessary or unavoidable, duplication must always be reported transparently, be properly attributed, and compliant with copyright requirements. If a manuscript contains text that has been published elsewhere, authors can avoid later problems by highlighting this to the Journal on submission.

7 Article categories

Reviews, Minireviews, Practice guidelines, Original articles, History of Asian medicine, Clinical (case) reports, Clinical vignettes, Technical reports, Brief communications, Controversies in medicine, Commentary, New developments, Clinico-pathological conferences, and Letters to the editor will be considered for publication.

7.1 Reviews

Asian biomedicine aims to provide narrative reviews as background for those in training, including graduate students and house officers or residents, and for the continuing education of practitioners and researchers, and others who wish an overview of the current status of a field. Authors are encouraged to submit reviews of various topics of relevance to Asia. Reviews should describe the current state of knowledge or practice, integrating recent advances with accepted principles and practice, or summarizing and analyzing the consensus view of controversial issues in knowledge or practice. Reviews should aim to be critical and to provide the reader with a balanced overview of an important and topical subject in biomedicine or health sciences, emphasizing factors such as cause, diagnosis, prognosis, therapy or prevention. They should cover aspects of a topic for which scientific consensus exists, and aspects that remain controversial and are the subject of ongoing scientific research. All articles and data sources reviewed should include information about the specific type of study or analysis, population, intervention, exposure, and tests or outcomes.

We would like reviews to be an original synthesis of ideas. The preparation of a review must, by its very nature, rely heavily on the ideas, observations, and reports of others. Therefore, it is important that the authors have exercised care in citing and quoting other articles. This precaution applies also to the use of one’s own published writing. The following guidelines are intended primarily to protect our authors from inadvertent infringements of copyright, any appearance of plagiarism, or accidental bias in assembling bibliographies.

Extra vigilance is required of literature review authors, for whom the task of materials assembly (today often a software cut-and-paste operation) must be separated carefully from those of information synthesis and fresh expression.
Ideally we would like the narrative review to define the current state of scientific knowledge, authors should strive to be fair, yet discriminating, in their selection of references. Including only those papers considered to be genuinely important; and not to clutter the bibliography with citations of marginal relevance to their topic merely for the sake of “completeness.” Without being so sparing with their references that they might appear to have minimized or disregarded the work of their competitors or newcomers to the field. Narrative reviews are based largely on unstructured expert opinion and should be identified as such; authors are encouraged to express their opinion rather than simply produce a review of the field with statements that are too general or miss important points. Authors should ensure that their chosen position is tenable. Guidance is available from the following articles among others: Gasparian AY, Ayvazyan L, Blackmore H, Kitas GD. Writing a narrative biomedical review: considerations for authors, peer reviewers, and editors. Rheumatol Int. 2011; 31:1409–17; Squires BP. Biomedical review articles: what editors want from authors and peer reviewers. CMAJ. 1989; 141:195–7; Green BN, Johnson CD, Adams A. Writing narrative literature reviews for peer-reviewed journals: secrets of the trade. J Chiropr Med. 2006; 5:101–17.

By contrast, systematic reviews include articles or data sources selected systematically and critically evaluated. Systematic reviews use explicit and reproducible criteria to assemble, appraise, and combine articles with a minimum of bias. Meta-analysis builds upon systematic review by using statistical techniques to pool and summarize data. Please follow Moher, Liberati, Tetzlaff, Altman, The PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: The PRISMA Statement, and its extensions as appropriate as detailed on the EQUATOR network (www.equator-network.org/home/) when reporting these types of reviews.

A review should include an abstract of not more than 250 words to inform readers of the authors’ reasons for reviewing the topic, their methods, their findings and their conclusions. Review articles are usually submitted by invitation only. However, unsolicited review articles will be given due consideration. Typical length: 6,000–10,000 words excluding references, Tables, and Figures, and no more than 100 references. Attention should be focused on articles published in the previous 5 years.

7.2 Minireviews

Review of special topics. Usually a narrative review concentrating on advancements in the field in the previous 5 years. Typical length: not more than 4,000 words excluding references, Tables, and Figures, and no more than 40 references.

7.3 Practice guidelines

Dealing with accepted medical, scientific, and social problem resolution. Clinical practice guidelines should follow the AGREE or RIGHT reporting tools for practice guidelines in health care, found at the EQUATOR Network (www.equator-network.org/home/). Typical length: not more than 10,000 words including abstract, excluding references, Tables, and Figures, and not more than 100 references.

7.4 Original articles

Original articles are full-length reports of current research which represent new and significant contributions to the field. We strongly recommend that authors refer to the minimum reporting guidelines for health research hosted by the EQUATOR Network (www.equator-network.org/home/) as indicated for various types of articles in parentheses as follows, and the FAIRsharing portal (https://fairsharing.org/) for reporting checklists for biological and biomedical research, where applicable, when preparing their manuscript. Original articles typically include clinical trials and intervention studies (CONSORT), studies of screening and diagnostic tests (STARD), laboratory (FAIRsharing) and animal studies (ARRIVE), cohort, case-control, and cross-sectional observational studies in epidemiology (STROBE), multivariable prediction models for individual prognosis or diagnosis (TRIPOD), biospecimen reporting (BRISQ), reliability and agreement studies (GRRAS), cost-effectiveness analyses (CHEERS), and surveys with high response rates (see STROBE-RDS for respondent-driven sampling studies, and Kelley, Clark, Brown, Sitzia. Good practice in the conduct and reporting of survey research. Int J Qual Health Care. 2003; 15:261–266). Authors should adhere to these guidelines when drafting their manuscript, and peer reviewers will be asked to refer to these checklists when evaluating such studies. Authors are expected to submit the checklist that is most appropriate for their manuscript type. If none of the guidelines listed above are suitable for the manuscript, the author is requested to either search for the most relevant set of guidelines supplied by the EQUATOR Network or explain during the submission process why none of the guidelines are appropriate for their study type.

Section headings based on IMRAD structure should usually be used: Abstract, Introduction, Materials and methods, Results, Discussion, Conclusion, Author contributions,
Conflicts of Interest Statement, Acknowledgments, and References.

The Introduction should provide a brief context or background to the subject of the manuscript, explain the importance or significance of the study, and clearly state the specific question, research objectives or purpose of, or hypothesis tested by, the study or observation. Avoid a detailed review of the literature in this section, but include sufficient background to understand the motivation for the study, and explain the experimental approach. Cite only directly pertinent references, and do not include data or conclusions from the work being reported.

The Methods section should describe the study design and methods. Describe the research methodology in sufficient detail so that others could duplicate the work. This section should specify the name of the animal (IACUC) or human ethics institutional review board (IRB) approved the research protocols appropriate and the approval reference number as appropriate (or that the research was exempt from approval and why), and if appropriate that the included participants provided documented informed consent. Manuscripts dealing with human studies should also declare that the study followed the principles of the contemporary revision of the Declaration of Helsinki. Manuscripts regarding animal studies should indicate that relevant national, international or local regulations were followed, and these should at least meet the standards of the National Institutes of Health Guide for the Care and Use of Laboratory Animals (Washington, D.C.: National Academy Press; 1996). All participants in human studies should provide documented informed consent and this information should be expressed in the manuscript. Documented informed consent including signed informed consent forms, witnessed and documented verbal consent, or tape recordings should be archived by the authors. The authors have to provide a written statement that they have received and archived documented informed consent from all patients, as required (see section 3). Copies of the letters of IRB or IACUC approval should be submitted with the manuscript and are required for acceptance. In all cases, the complete name of the IRB or IACUC, preferably with the approval number should be provided in the manuscript. Include the study setting and dates, patients/participants with inclusion and exclusion criteria, relevant demographic details, patient samples or animal specimens used, the essential features of any interventions, the main outcome measures, the laboratory methods followed, or data sources and how these were selected for the study, instrumentation and key reagents, and state the statistical procedures employed in the research. Specify the statistical software package(s) and versions used. The symbol ®, or letters ™ or SM, should not be used in scientific journal articles, but to comply with trademark law (U.S. Federal Trademark Dilution Act of 1996. Pub No. 104–98, 109 Stat 985) the initial letter of a trademarked word should be capitalized. The manufacturer of any drug, reagent, or equipment mentioned in the manuscript should be mentioned, with a catalog number as appropriate, such as for antibodies, enzymes, probes, or specialist reagents. Location details can usually be omitted because such information is now so freely available on the Internet that we consider it is no longer necessary to continue to require this, as consistent with the American Medical Association house style since October 4, 2011. We recommend that commercial suppliers of reagents or instrumentation be identified only when the source is critical to the outcome of the experiments. For example, it is usually not necessary to specify the source of common reagents such as buffer salts. However, do use a general statement such as “all chemicals used were analytical grade or higher” as appropriate. Nonproprietary names or descriptive phrasing is preferred to proprietary names for devices, equipment, and reagents, particularly in the context of general statements and interchangeable items (e.g., urinary catheters, intravenous catheters, pumps). If the use of proprietary names is necessary for clarity or to replicate the study, proprietary names should be given at first mention along with the nonproprietary name. In such cases information regarding the manufacturer or supplier (and sometimes location, but see above) is important, and authors should include this information in parentheses after the name or description. It is not necessary to identify the legal status suffix of the supplier, e.g. Ltd., Inc., GmbH, AG, and S.A., unless this forms part of the company name and common usage is preferred (e.g. DuPont for the E. I. du Pont de Nemours and Company).

On occasion, a trademark owner will request that its trademark or trade name appear in all capital letters or a combination of capital and lowercase letters often with the trademark symbol. Authors and Editors are not required by law to follow such requests. It is preferable to use an initial capital letter followed by all lowercase letters (e.g., Xerox, Kodak) unless the trademark name is an abbreviation (e.g., IBM, JAMA) or uses an intercapped construction (e.g. PubMed, iTunes). Online databases, if trademarked, can be listed in all capital letters (e.g., MEDLINE, CINAHL). Use only an initial capital letter, not all capitals, for devices, equipment mentioned in the manuscript should be mentioned, or interchangeables (e.g., urinary catheters, intravenous catheters, pumps). If the use of proprietary names is necessary for clarity or to replicate the study, proprietary names should be given at first mention along with the nonproprietary name. In such cases, information regarding the manufacturer or supplier (and sometimes location, but see above) is important, and authors should include this information in parentheses after the name or description. It is not necessary to identify the legal status suffix of the supplier, e.g. Ltd., Inc., GmbH, AG, and S.A., unless this forms part of the company name and common usage is preferred (e.g. DuPont for the E. I. du Pont de Nemours and Company).

The Results section should comprise the study results presented in a logical sequence, supplemented by Tables or Figures, or both, giving the main or most important findings first. Take care that the text does not repeat data that are presented in Tables or Figures. Emphasize and summarize only the essential features of the main outcome measures, and the
main results. Where appropriate, findings should be quantified and presented using appropriate indicators of measurement error or uncertainty (e.g., confidence intervals). Data should be presented with an appropriate degree of precision and statistical data should be provided for all tested differences. See section 8.4.13.

The Discussion should be used to emphasize the new and important aspects of the study, placing the results in context with the entire published literature, the implications of the findings, and the conclusions that follow from the study results. Begin with a description of what your study found in relation to the purpose or objectives as stated in the Introduction. To avoid redundancy and for conciseness, do not repeat in detail data or other information given in other parts of the manuscript, such as in the Introduction or the Results section. Describe rather than repeat results given earlier. Although some degree of speculation as to the importance of the observations is permissible if identified as such, avoid making conclusions unrelated to the data presented. For experimental studies, it is useful to begin the discussion by briefly summarizing the main findings, then explore possible mechanisms or explanations for these findings, compare and contrast the results with other relevant studies, clearly state the limitations of the study, and explore the implications of the findings for future research and for clinical practice. Link the conclusions with the goals of the study, but avoid overstating them by making unqualified statements, and presenting conclusions not adequately supported by the data. In particular, distinguish between clinical importance and statistical significance, and avoid making statements on economic benefits and costs unless the manuscript includes the appropriate economic data and analyses. Please avoid unsupported claims of primacy or priority purporting that your study is the first (or largest of its type) or alluding to work that has not been completed. For example, avoid statements such as, “This paper is the first to report...” because even the most thorough searches may fail to reveal all instances of similar work. Although topics that require future research can be mentioned, it is unnecessary to state that “further research is needed.” State new hypotheses when warranted, but clearly identify them as such. A final summary is not necessary, because this information should be provided in the Abstract and the first paragraph of the Discussion. Instead provide Conclusions that can specifically be drawn from the current work, do not overstate your results or make conclusions related to other studies.

We prefer that manuscripts are organized with the Results and Discussion sections separated followed by a Conclusion as detailed above. However, exceptions for combined Results and Discussion sections can be made if this is justified in additional comments you would like to send to the publication office during the submission process in Editorial Manager.

Typical length: 3,000–5,000 words excluding references, Tables, and Figures, and not more than 50 references.

7.5 History of Asian biomedicine

Articles concerning the history and development of biomedicine in the Asian region. Includes articles regarding early medical efforts, specific milestones, and biographies of biomedical pioneers and past biomedical “giants” associated with the Asian region. All contributions should be thoughtful, and documented by the proper citation of original works or secondary sources. Typical length: not more than 4,000 words excluding references, Tables, and Figures, and not more than 50 references.

7.6 Clinical reports

Clinical reports include case reports. We want to publish cases with clinical valuable lessons and documenting the state of current practice. Therefore, common cases that present a diagnostic, ethical or management challenge, or that highlight aspects of mechanisms of injury, pharmacology or histopathology are deemed to be of educational value. In addition, we encourage reports of global health cases and medicine practiced in Asian settings. We want to publish cases worthy of discussion particularly around aspects of differential diagnosis, decision making, management, clinical guidelines, and pathology. Cases will be judged on clinical interest and educational value, not necessarily novelty or rarity. Please follow CARE reporting guidelines available from the EQUATOR network and https://www.care-statement.org/.

You must have documented informed consent from patients (or parents, or legally authorized guardians or representatives in the case of minors or incapacitated adults), whether identifiable or not, before submitting case reports or case series to Asian Biomedicine. Please anonymize the patient’s details as much as possible, e.g., specific ages, ethnicity, and occupations without omitting details essential to the clinical message or scientific purpose of the article. We will not send your manuscript for review without explicit consent from the patient, or legally authorized guardian or representative on our Patient privacy and confidentiality form available at the home (welcome) page of Editorial Manager for Asian Biomedicine (please see section 5 for more details). If the patient is dead, the authors must seek permission from a relative (ideally the next of kin). If you do not have signed
consent from a deceased patient, guardian or family, the head of your medical team/hospital or legal team must take responsibility that exhaustive attempts have been made to contact the family and that the paper has been sufficiently anonymized so as not to cause harm to the patient or their family. You will need to provide a signed document to this effect. Typical length: not more than 6,000 words excluding references, Tables, and Figures, and not more than 40 references (See also section 7.7).

7.7 Clinical vignettes

A clinical vignette is a short report of one or more patient-related cases or technical issues, and scenarios that have educational value for a wider audience, for example by illustrating a new disease entity or a prominent or unusual clinical feature of an established disease, highlighting an area of clinical controversy, a technical improvement, or illustrating a unique patient safety issue.

Clinical vignettes should be based on patients for whom at least one of the author(s) had cared during the course of the patient’s illness. Clinical vignettes should describe clinical conditions that illustrate unique or important teaching points; provide insight into clinical practice, education, or research in either outpatient or hospital settings; illustrate important clinical problems commonly encountered by internists, such as diagnostic, therapeutic, or management dilemmas, including those complicated by factors such as low health literacy or language barriers. Practical or unique technical procedures or instrumentation may also be appropriate. Clinical vignettes should include a discussion of relevant literature, but they are not intended for presentation of scientific or research data.

Consider presenting a case if it increases awareness of a condition, suggests a proper diagnostic strategy, or demonstrates a more cost-effective approach to management. Alternatively, a case can be presented because it represents an unusual presentation of a relatively common condition. Other aspects include an unusual complication of a disease and its management.

Submissions should conform to the following organization: 1. A short Introduction typically describing the context of the case, and explaining its relevance and importance. 2. Learning objectives stating what the practitioners should be able to do with the case information. 3. Case description: briefly summarize the case presentation. Include a history of the present illness, pertinent positives, negatives, key past medical history/social history (personal habits, living situation, job)/family history/medications, key physical exam, diagnostic studies, interventions. Highlight key elements from clinical course. Be complete without obscuring the essence of the case with irrelevant details. 3. Discussion: a concise discussion, highlighting important facts or teaching points gleaned from the case and/or review of the literature. Please also follow the CARE reporting guidelines available from the EQUATOR network and https://www.care-statement.org/. Please avoid claims that your case is the “first” to describe a particular phenomenon (see section 7.4).

You must have documented informed consent from patients (or parents or legally authorized guardians or representatives in the case of minors or incapacitated adults) whether identifiable or not before submitting case reports to Asian Biomedicine. Please see section 7.6 for further details). Typical length: not more than 2,000 words excluding references, Tables, and Figures, and not more than 20 references.

7.8 Technical report

Technical reports are detailed reports of novel techniques. Typical length: not more than 6,000 words excluding references, Tables, and Figures, and not more than 40 references.

7.9 Brief communication

Brief communications are short research articles. Please see the requirements for Original articles (section 7.4), and note that we strongly recommend that authors refer to the minimum reporting guidelines for health research hosted by the EQUATOR Network as indicated for various types of articles in parentheses as follows, and the FAIRsharing portal (https://fairsharing.org/) for reporting checklists for biological and biomedical research, where applicable, when preparing their manuscript. Peer reviewers will be asked to refer to these checklists when evaluating such studies. Authors are expected to submit the checklist that is most appropriate for their manuscript type. Typical length: not more than 3,000 words, excluding references, Tables, and Figures, and not more than 20 references.

7.10 Controversies in biomedicine

Articles concerning various controversial medical topics. Two authors should write on opposing sides of an issue of relevance to the Asian region. Each weighing the pros and cons of their argument. A brief introductory paragraph should be included to explain what the controversial issue is and what the two arguable sides are. Typical length: not more than
3,000 words excluding references, Tables, and Figures, and not more than 20 references.

7.11 Commentary

Comments dealing with scientific and social issues. Typical length: not more than 3,000 words, excluding references, Tables, and Figures, and not more than 20 references.

7.12 New developments

Bioscience developments of interest to the Asian region. Summaries of important publications and books from Asia. Summaries of doctoral theses from the region.

7.13 Clinicopathological conferences

Clinicopathological conferences (CPCs) emphasize new ideas in medical science that shed light on the approach to diagnosis, disease management, or an emerging aspect of pathophysiology. The editors welcome enquiries about cases that might be appropriate subjects for CPCs, and invite submissions of CPCs. No abstract is necessary. On the second page, begin the case presentation using the following headings (as applicable): Case presentation (history of the present illness, past medical history, social and family history, review of systems, physical examination, laboratory evaluation), Case Summary, Differential diagnosis, Discussion, the Patients' course, Final diagnosis. CPCs may be published twice per year at the discretion of the Editors. You must have signed informed consent from patients (or parents, or legally authorized guardian or representative in the case of minors or incapacitated adults) before submitting to Asian Biomedicine, for details see section 7.6). Typical length: not more than 6,000 words, excluding references, Tables, and Figures, and not more than 50 references.

7.14 Letters to the Editor

Letters are welcome in response to previously published articles in Asian Biomedicine, or other brief technical or clinical notes of general interest. Letters should have a title, no more than 4 authors, include appropriate references and the corresponding author’s mailing and e-mail addresses. Letters are edited, sometimes extensively, to sharpen their focus. They may be sent for peer review at the discretion of the Editors. A copyright transfer form must be signed. Letters are selected based on clarity, significance, and space. Typical length: not more than 1000 words excluding references, 1 Table or Figure, and not more than 5 references.

8 Manuscript preparation

Text should be presented double-spaced A4-sized (297 mm × 210 mm) pages, with outer margins of 2.5 cm and left aligned text without justification. A manuscript should include a title page, abstract, text, references, conflicts of interest statement, acknowledgments, author contributions, and Figures and Tables as appropriate. Pages should be numbered consecutively, beginning with the title page. Please submit MS Word files through the Editorial Manager system (see sections 1.2 and 7).

8.1 Title page

The title page (see section 1.2) should contain the following information:
• article category
• article title (see section 8.2)
• author names (full first name, and middle names or initials (optional), family name last) and the institutions with which they are affiliated; indicate all affiliations with a superscripted number (to avoid confusion as being part of the author’s name) after the author’s name and in front of the matching affiliation. If an author’s affiliation has changed during the course of the work, the author should list the affiliation at the time that the research (or most significant portion of the research) was conducted. Their current affiliation may be given in the Acknowledgments section, where the change of affiliation can be explained if considered necessary. The corresponding author should use their current address for correspondence, but their affiliation at the time when the work was conducted. The use of an ORCID is encouraged. See https://orcid.org/. In the published article, at least one, or optionally more than one author must be designated with an asterisk as the author(s) to whom reader correspondence regarding the published article may be addressed. This does not need to be the same as the corresponding author, but ideally should be somebody whose contact details are not likely to change in the near future, and must except in unusual circumstances have a professional or institutional email address. We appreciate those in supervisory or administrative roles may be more senior authors who may not have the time to communicate with journal editors, and other authors. They may also not have sufficient time, experience or
patience for the online submission processes, but may be
the most effective author to whom reader correspondence
regarding the published article may be addressed.
• short title not exceeding 50 characters.

8.2 Article title
The title should be informative and provide a concise
description of the complete article and include information that, along
with the Abstract, will make electronic retrieval of the article
sensitive and specific. Please include information about the
study design as a part of the title (particularly important for
randomized trials, systematic reviews, and meta-analyses).
Asian Biomedicine additionally requires a short title, not more
than 50 characters (including letters and spaces) on the title
tape (to be used as a running head).

8.3 Abstract and Keywords
Please provide an Abstract (of no more than 250 words for
brevity and to avoid automatic truncation by electronic indexing
systems) structured as appropriate under 5 headings: Background
(context for the study), Objectives (study’s purpose or
aims; please avoid using verbs such as explore or examine in
the objectives because it is later difficult to determine if the
investigators were successful), Methods (basic procedures
(selection of study participants, settings, measurements, analytical methods)), Results (main findings (giving specific
effect sizes and their statistical and clinical significance,
if possible)), and Conclusions. For Clinical reports please
limit the abstract to 150 words. An unstructured abstract may
be used for Reviews, History of Asian Medicine, Controversies,
and Commentaries. No abstract is required for Clinical
vignettes, CPCs, New developments, or Letters to the Editor.
Please avoid the use of abbreviations in the Abstract unless
they are absolutely necessary and used more than once. Please
do not cite references in the abstract unless provided in full
as part of the word count. We can accept Abstracts up for 300
words in submissions for revision to our 250-word limit at the
discretion of the editors.
Clinical trial abstracts should include items that
the CONSORT group has identified as essential (www.
consort-statement.org/resources/downloads/extensions/
consort-extension-for-abstracts-2008pdf/) including the clinical
trial registration number at the end of the abstract.
Keywords. Please provide from 5 to 10 key words or
short phrases in alphabetical order. The use of keywords
taken from the Medical Subject Headings (MeSH) list of the
U.S. National Library of Medicine at the following URL is
couraged (http://www.nlm.nih.gov/mesh/meshhome.html)
and will facilitate searching and retrieval of published articles.

8.4 Main Text
The text for most articles should be organized into the fol-
lowing sections: Introduction, Materials and methods, Results,
Discussion (IMRAD), Conclusion, Author contributions, Con-
flicts of interest, Acknowledgments, and References. For Clin-
cial reports Patients and methods might be used as appropri-
ate. See individual article types in section 7 for further details.
Please refer to the American Medical Association Manual of
Style (9th edition or later) for matters of style.

8.4.1 Abbreviations
Asian Biomedicine discourages the use of abbreviations and
discourages the use of abbreviations and acronyms, and asks that authors avoid the use of abbreviations
in the abstract and title entirely unless they are absolutely
necessary and used more than once. Abbreviations impose a
burden on a reader because the reader must first decipher the
author’s code. This task distracts the reader from concentrat-
ing on the content of an article. A reader should not need a spe-
cialized cipher key to understand an article. An article should
not be “alphabet soup”. It is acceptable to substitute a standard
abbreviation for an unwieldy word or phrase appearing more
than, say 5 times in a manuscript. An abbreviation should
never replace one short word. A simple way of avoiding abbre-
viation is to use a substitute word. For example, instead of
writing “IRL” for “inspiratory resistive load”, simply write
“load” or “the load” after first stating what type. This will
facilitate the “flow” of the text.
Please limit the use of nonstandard abbreviations. Where
a term/definition will be continually referred to, it must be
written in full when it first appears in the text, followed by the
abbreviation in parentheses immediately after it. Any abbrevi-
atations should be defined at first mention in the Abstract, text,
and in each Table and Figure (which must be understood on
its own without reference to the main text). Thereafter, the
abbreviation should be used consistently. Avoid first defining
abbreviations in any section heading; if an abbreviation has
previously been defined in the text, then the abbreviation may
be used in a subsequent section heading. Restrict the number
of abbreviations to those that are absolutely necessary. For lists
of standard abbreviations please consult the Council of Biology
Editors Style Guide or other standard sources such as the Amer-
ican Medical Association Manual of Style (9th edition or later).
Avoid using terms like Group 1 or Group A; readers should not have to remember what Group 1 or Group A stands for. Instead, write the “treated patients” or the “control group” or something else as appropriate.

8.4.2 Jargon, slang, colloquialisms, idioms, homonyms, euphemisms, and clichés

Specialized medical jargon and medical slang should be avoided. Colloquialisms, idioms, and vulgarisms should also be avoided in formal scientific writing. “When these words appear in medical manuscripts or in medical conversation, they are unintelligible to other scientists, particularly those of foreign countries; they are not translatable…” (Words and phrases. In: Fishbein. Medical writing: the technic and the art. Chicago: American Medical Association; 1938:46p).

Carefully check homonyms that sound alike, but are spelled differently and have different meanings. Euphemisms are indirect terms used to express something unpleasant. Directness is better in scientific writing (“The patient died.”). Avoid “sacrificed” when referring to animals (note carcass not cadaver); use “killed”, or better “humanely killed” instead. Avoid clichés “like the plague”.

Please find more detailed discussion of correct and preferred language and grammar in the American Medical Association Manual of Style (9th edition or later).

8.4.3 Numbers

Numbers that begin a sentence should be spelled out as words. The use of “one” in running text or other numbers, zero or two should usually be written as words where use of a numeral would place unintended emphasis on a precise quantity or would be confusing. Laboratory parameters, and all quantities with units of measure, time, temperature, number of subjects, length, area, mass, and volume should be expressed using arabic numerals. Please use an en rule (–) and not a hyphen (-) to indicate spans of numbers, e.g. patients aged 6–64 years, and minus signs.

8.4.4 Units

In general, Système International (SI) units must be used (updated (2014) 8th edition of the SI Brochure, which defines and presents the Système International d’Unités, the SI (known in English as the International System of Units is available at http://www.bipm.org/en/publications/si-brochure/). However, see the AMA Manual of Style for guidance on use and exceptions, e.g., blood pressure values, which should be reported in mmHg, temperatures to be given in degrees Celsius (°C), and enzyme activity that may be reported in International Units. Other non-SI units used informally in health care and biomedical fields may be used at the discretion of the editors, but may require definition. Please use the metric system for the expression of length, area, mass, and volume. Please use a space between figures and the symbols (e.g. 25 kg). The degree symbol is an exception, written without spaces (e.g. 12°, and 37°C consistent with the AMA Manual of Style. Please use a degree symbol and not a superscripted “°”. The correct abbreviations of the hours and seconds are “h” and “s” respectively and not “hr” and “sec.” The symbol used for liter should be an uppercase “L” as consistent with the AMA Manual of Style and the U.S. National Institute of Standards and Technology (NIST) recommendations.

8.4.5 Names of drugs, devices, and other products

Use the recommended International Nonproprietary Name (rINN) for medicinal substances, unless the specific trade name of a drug is directly relevant to the discussion. Generic or chemical drug names should appear in lowercase letters in the text and not be abbreviated. If a specific proprietary drug needs to be identified, the copyright or brand name with an initial capital may appear only once in the manuscript in parentheses following the generic or chemical name the first time the drug is mentioned in the text. The use of trademark names should not be used with trademark symbols, e.g. ® or ™, but the trademarked word should have an initial capital (see section 7.4). For devices and other products (chemicals, reagents, or equipment), the specific brand or trade name, and the manufacturer should be provided the first time the device or product is mentioned in the text. Thereafter, the generic term (if appropriate) should be used.

When the administration of drugs is described, intra-articular, intracardiac, intramuscular, intrathecal, intravenous, intraventricular, intravitreal, oral, parenteral, rectal, subconjunctival, subcutaneous, sublingual, topical, and transdermal are acceptable terms when these are the usual or intended routes of administration.

8.4.6 Gene nomenclature

Current standard international nomenclature for genes should be adhered to. For human genes, use genetic notation and symbols approved by the HUGO Gene Nomenclature Committee (http://www.genenames.org). You may also refer to the
resources available on PubMed at http://www.ncbi.nlm.nih.gov/guide/genes-expression. The Human Genome Variation Society has a useful site that provides guidance in naming mutations at http://www.hgvs.org/mutnomen/index.html. In your manuscript, genes should be typed in italic typeface, and include the accession number at first mention if appropriate.

8.4.7 Genomic and related data

Where there is a community established norm for data sharing, authors are required to deposit their genomic or other relevant datasets into publicly available databases on the date of publication. For the purposes of evaluating the manuscript, authors should provide the editors and reviewers with access to the datasets when the manuscript is submitted to the Asian Biomedicine; information about accessing the datasets can be included in the Materials and methods section. A data accessibility statement may be required.

1. Microarray data: it is the authors’ responsibility to ensure that all data collected and analyzed in their experiments adhere to the Minimal Information About A Microarray Experiment (MIAME) guidelines. Review the MIAME checklist at the following URL: http://www.mged.org/Workgroups/MIAME/miame.html

Appropriate public databases include, but are not limited to: ArrayExpress, Gene Expression Omnibus (GEO), or CIBEX.

2. Nucleotide and protein sequences: DNA Sequences, RNA Sequences, and Protein Sequences. Sequences should be deposited according to Minimum information about any (x) sequence (MiXS) guidelines available at http://www.ebi.ac.uk/ena/submit/mixs-checklists.

Appropriate public databases include, but are not limited to: European Molecular Biology Laboratory (EMBL/EBI), GenBank, or DNA Data Bank of Japan (DDBJ), Uniprot, NCBI, dbVar (genetic polymorphisms) and ClinVar.

3. Polymerase chain reaction (PCR): a set of guidelines covering key parameters of every qPCR assay that are essential for allowing reviewers, editors and readers to evaluate the technical merits of scientific publications using qPCR technology are found in the MIQE précis: practical implementation of minimum standard guidelines for fluorescence based quantitative real-time PCR experiments (see Bustin et al. BMC Molecular Biology 2010; 11:74. doi: 10.1186/1471-2199-11-74).

4. Crystallographic data for small molecules: Manuscripts reporting new 3-dimensional structures of small molecules from crystallographic analysis should include a .cif file and a structural figure with probability ellipsoids checked using the IUCR’s CheckCIF routine (http://checkcif.iucr.org/), and a PDF copy of the output must be included with the submission, together with a justification for any alerts reported. Crystallographic data for small molecules should be submitted to the Cambridge Structural Database (https://www.ccdc.cam.ac.uk/) and the deposition number referenced appropriately in the manuscript. Full access must be provided on publication.

8.4.8 Plant (or other) extracts

Asian Biomedicine can no longer accept data for crude extracts that have not been characterized by analysis of their major constituents (e.g. by HPLC, NMR, and mass spectroscopy (MS)). Identification of extracts should be supported by at least an HPLC or gas chromatography (GC) trace, which should be included with the manuscript. Extraction and isolation of compounds should be described in detail. The kind and amount of material, solvents, and extraction methods must be indicated. Please consider standardizing your various extracts by means of HPLC, HPLC-MS, or HPLC-NMR—fingerprinting inclusive of the identification and quantitation of the typical constituents, secondary metabolites, and main bioactive compounds, which are, or might be responsible for the pharmacological effects or activity studied. HPLC or GC followed by a suitable detection method, or by enzyme-linked immunosorbent assays, radioimmunoassays, ligand binding assays, various sensors, or fluorometric assays, to name a few, might be appropriate. The methods should be described in detail: including apparatus, columns, solvent systems, gradient, flow rate, and detection. If you do not possess the required analytical equipment or expertise, please consider seeking cooperation with a phytochemical laboratory. Without standardization of the plant (or other) extracts, the results presented cannot be pharmacologically reproduced, and we cannot accept them for publication. We would like to see evidence at a molecular level supportive of any speculation regarding the mechanism of action of putatively active components, e.g. levels of components (that might be responsible for the activity). This allows a better comparison against positive controls that might include putatively active components. A negative control is also necessary. It is usually desirable to demonstrate a dose–activity dependence. This would require at least 3 dose–responses.

Collecting or research on plants or other specimens (either cultivated or wild) from botanical gardens, national parks, or research forests and the like, must comply with institutional, national, or international guidelines and this needs to
be explicitly reported in the Methods section of manuscripts. Some statement should be made that the collection was approved, did not require any specific permits (which should be specified if required), and did not violate any endangered or protected species. The requirement for specific permits may be waived for field studies in locations are not privately owned or protected in any way. The month and year when the plants were collected should be stated, and ideally, the exact collection location should be provided using a GPS navigation tool. We require that authors comply with the Convention on Biological Diversity and the Convention on the Trade in Endangered Species of Wild Fauna and Flora (see section 3).

Investigators who purchase dried “herbal remedies” or other materials from companies need to specify the name of the supplier and their location and ideally deposit samples in an herbarium.

Manuscripts must report the presence of specimen plants in a specified major regional herbarium, for access by future workers, and include the reference number of the voucher specimen of the plant specimen. The part of the plant used and the full name, highest academic degree, and affiliation of the expert botanist who examined it should be reported. Please provide complete information for the correct nomenclature, the scientific name (in italics), the author of this name and the family under the heading “Plant Material.” If this is in doubt, the Index Kewensis (electronic Plant Information Centre ePIC, Royal Botanic Gardens, Kew, UK: http://www.kew.org/epic), and/or the International Code of Botanical Nomenclature (www.bgbm.fu-berlin.de/iapt/nomenclature/code/tokyo-e/default.htm) should be followed. It is also helpful to mention the commonly used name for the plant that is the focus of your study.

For Chinese or ethnic traditional medicine (e.g. Kampo, Ayurvedic, or Korean traditional medicine) please use a recognized source for plant names. Herbal names in traditional Chinese medicine have been a problem in terms of capitalization and italicization, because using italicization can get names confused with formal Linnaean classification. A helpful and authoritative online resource is from the WHO: http://apps.who.int/medicinedocs/en/d/Js2200e/20.html (and related pages). Please use common herb names are in plain text and the formal species names in italics. For example, Radix Glycyrrhizae consists of the dried roots and rhizomes of Glycyrrhiza glabra L. and its varieties (1–7) or of Glycyrrhiza uralensis Fisch. (6, 7) (Fabaceae). Other useful sources are Chan et al. Good practice in reviewing and publishing studies on herbal medicine, with special emphasis on traditional Chinese medicine and Chinese Materia Medica. Journal of Ethnopharmacology 2012; 140:469–75 and Ho-Dzun, Knüpffer, and Hammer. Additional notes to the checklist of Korean cultivated plants (5). Consolidated summary and indexes. Genetic Resources and Crop Evolution 1997; 44:349–91.

The structural analysis of new plant substances is now so routine that papers reporting a single novel compound of expected structure (e.g. a new triterpene fatty acid ester) are rarely acceptable, unless other novel information regarding the plant is included.

Manuscripts describing routine bioactivity screening (e.g., antimicrobial, antioxidant, antifeedant, etc.) of crude extracts or predictable and/or unexceptional bioactivity (e.g., antioxidant activity of polyphenolics, antimicrobial activity of essential oils), are generally not acceptable.

Uncritical ethnopharmacological investigations, where a list of plants and their use are simply recorded will not be accepted.

8.4.9 Cell lines

If human cell lines are used, authors are strongly encouraged to include the following information in their manuscript:

• The source of the cell line, including when and from where it was obtained.
• Whether the cell line has recently been authenticated and by what method. Cell line authentication is recommended; e.g., by karyotyping, isozyme analysis, or short tandem repeats (STR) analysis—and may be required during peer review or before publication.
• Whether the cell line has recently been tested for mycoplasma contamination.
• For established cell lines, the Methods section should include: a reference to the published article that first described the cell line; AND/OR the cell line repository or company the cell line was obtained from, the catalogue number, and whether the cell line was obtained directly from the repository/company or from another laboratory.
• For de novo (new) cell lines, including those given to the investigators as a gift, authors must follow our policies for research involving human subjects or animals, as appropriate. The ethics statement must include: details of institutional review board or ethics committee approval; AND for human cells, confirmation of written informed consent from the donor, guardian, or next of kin.

Further information is available from the International Cell Line Authentication Committee at: https://standards.atcc.org/kwspub/home/the_international_cell_line_authentica-tion_committee-iclac/_ We recommend that authors check the NCBI database for misidentification and contamination

8.4.10 Blots and gels

Manuscripts reporting results from blots (including western blots) and electrophoretic gels should follow the following guidelines:

- In accordance with our policy on image manipulation (see section 8.10), the image should not be adjusted in any way that could affect the scientific information displayed, e.g. by modifying the background or contrast.
- All blots and gels that support results reported in the manuscript should be provided.
- Original uncropped and unadjusted blots and gels, including molecular size markers, should be provided in the Figures.
- Lanes should not be overcropped around the bands; the image should show most or all of the blot or gel.
- Any nonspecific bands should be shown and an explanation of their nature should be given.
- The image should include all relevant controls, and controls should be run on the same blot or gel as the samples.
- A Figure panel should not include composite images of bands originating from different blots or gels. If the figure shows nonadjacent bands from the same blot or gel, this should be clearly denoted by vertical black lines and the figure legend should provide details of how the figure was made.

8.4.11 Antibodies

Manuscripts reporting experiments using antibodies should include the following information:

- The name of each antibody, a description of whether it is monoclonal or polyclonal, and the host species.
- The commercial supplier or source laboratory. The catalogue or clone number and, if known, the batch number.
- The antigen(s) used to raise the antibody.
- For established antibodies, a stable public identifier from the Antibody Registry.

The manuscript should also report the following experimental details.

- The final antibody concentration or dilution.
- A reference to the validation study if the antibody was previously validated. If not, provide details of how the authors validated the antibody for the applications and species used.

We encourage authors to consider adding information on new validations to a publicly available database such as Antibodypedia or CiteAb.

8.4.12 Tumor cell lines and anticancer activity

In manuscripts that present results of biological studies with tumor cell lines authors should pay special attention to the U.S. National Cancer Institute (NIH) guidelines for cancer drug discovery studies. Compounds that suppress the growth of, or kill, isolated tumor cell lines grown in culture should be referred to as either “cytostatic” or “cytotoxic”, as appropriate. The term “anticancer” should be reserved for compounds that show specific activity in human-based clinical studies (see Suffness M, Douros JJ. Nat. Prod. 1982; 45:1–14). Some flexibility in this system is afforded in the description of compounds that show activity in molecular-targeted antitumor assays.

Compounds should be compared against a suitable positive control substance and follow accepted guidelines when represented as “active.” For example, a cytotoxic pure substance when tested against a cancer cell line would exhibit an IC₅₀ value of <10 µM (or 4–5 µg/mL). Only sufficiently potent or exceptional activities will be considered. Report the biological activities by listing IC₅₀ values and minimal inhibitory concentrations (MICs) for the active substance being studied, and a positive control or reference material. A negative control should also be used. Results must be based on adequate statistics. Positive controls (reference/standard compounds) and at least 3 dose responses for conventional pharmacological experiments should be included.

8.4.13 Statistical requirements

Conclusions reported in Asian Biomedicine should be based on adequate statistics that incorporate appropriate tests of significance, account for the type of data distribution, and are based on the number of experimental observations required for the application of the respective statistical method.

Use correct nomenclature for statistical methods. Descriptive statistics should follow the scales used in data description. Inferential statistics are important for interpreting results and should be described in detail.

All P values should be presented to an appropriate degree of precision. In brief, P values should be expressed to 2 digits to the right of the decimal point (regardless of whether the P value is significant), unless P < 0.01, in which case the P value should be expressed to 3 digits to the right of the decimal point. (One exception to this rule is when rounding P from 3 digits to
2 digits would result in \( P \) appearing nonsignificant, such as \( P = 0.046 \). In this case, expressing the \( P \) value to 3 places may be preferred. The same holds true for rounding confidence intervals that are significant before rounding, but nonsignificant after rounding. Usually, with the exception of some genetics studies, the smallest \( P \) value that should be expressed is \( P < 0.001 \), because additional zeros do not convey useful information; the largest \( P \) value that should be expressed is \( P > 0.99 \). Please refer to the American Medical Association Manual of Style (9th edition or later) for a glossary of statistical terms and advice regarding significant digits and rounding numbers.

Percentages are usually inappropriate if the denominator is less than about 20. The cutpoint of 20 to indicate a small sample is reasonable, but arbitrary. Especially in small samples, percentages can be misleading because the size of the percentage can be so much greater than the number it represents. e.g. “In this experiment, 33% of the rats lived, 33% died, and the third got away.” (Lang and Secic. How to report statistics in medicine, 2nd ed. Philadelphia: American College of Physicians; 2006) Percentages may still be useful for comparison (38% vs 37% is an easier comparison than 5/13 vs 7/19). The actual numerator and denominator must be given if the denominator is less than about 20. Percentages are often unnecessary if the denominator is less than about 100, although they may be useful for ease of comparison of values with different denominators. Generally, when proportions or percentages derived from a mix of large (>100) and smaller (<100) denominators occur nearby, report all to the decimal level appropriate for the smallest denominator. Actual numbers should also be given. Percentages should not have a decimal place if the denominator is less than about 200; use integers (whole numbers). If the denominator is between 200 and 500, then one decimal place can be used, but is not required.

See Lang and Altman. Basic statistical reporting for articles published in biomedical journals: the “Statistical Analyses and Methods in the Published Literature” or the SAMPL Guidelines Int J Nurs Stud. 2015; 52:5–9 for further guidance. Reviewers will be asked to check the statistical methods, and the manuscript may be sent to our consultants for specialist statistical review if considered necessary.

### 8.4.14 Qualitative research

Qualitative research studies use nonquantitative methods to address a defined research question that may not be accessible by quantitative methods, such as people’s interpretations, experiences, and perspectives. The analysis methods are explicit, systematic, and reproducible, but the results do not involve numerical values or use statistics. Examples of qualitative data sources include, but are not limited to, interviews, text documents, audio/video recordings, and free-form answers to questionnaires and surveys. Qualitative research studies should be reported in accordance to the Consolidated criteria for reporting qualitative research (COREQ) checklist. Further reporting guidelines can be found in the EQUATOR Network’s Guidelines for reporting qualitative research.

### 8.4.15 Personal communications and unpublished data

These sources should not be included in the references list, but may be described in the text and should be referred to as “personal communications” and “unpublished observations.”

The author(s) must give the full name and highest academic degree of the person, the date of the communication, and indicate whether it was in oral or written (letter, fax, e-mail) form. A signed statement of permission should be included from each person identified as a source of information in a personal communication or as a source for unpublished data.

### 8.5 Authors’ contributions

The contributions of each author to the work that is reported should be clearly and briefly using their initials and stated according to ICMJE recommendations (see Section 1.3.2).

### 8.6 Conflict of interest statement

Please ensure that the initials of each author listed in your manuscript appears in the Conflicts of interest statement. If no authors have any potential conflict of interest, then it may be sufficient to state “The authors have each completed the International Committee of Medical Journal Editors Form for uniform Disclosure of Potential Conflicts of Interest. No authors have any potential conflict of interest to disclose/All authors have nothing to disclose/None of the authors disclose any potential conflict of interest”. See section 2.

### 8.7 Acknowledgments

General acknowledgments for consultations and statistical analysis should be listed concisely, including the names of the individuals who were directly involved. Consent should be obtained in writing from those individuals before their names are listed in this section (see section 2. Authorship
statement). All financial and material support for the research, work, writing and editorial assistance from internal or external agencies, including commercial companies, should be clearly and completely identified. Previous publication of the work as an abstract of a poster or oral presentation at a conference can be presented here. If a study is funded by an agency with a proprietary or financial interest in the outcome of the study, the corresponding author must include the following statement in the comments to the editors and in the Acknowledgments section upon submission: “I had full access to all of the data in this study and I take complete responsibility for the integrity of the data and the accuracy of the data analysis.” (see section 2). Changes in author affiliation and explanations if necessary can be presented here (see Section 8.1). Do not include thanks to anonymous referees or editors, or effusive comments.

Group authorship: if you would like the names of the individual members of a collaboration Group to be searchable through their individual PubMed records, please ensure that the title of the collaboration Group is included on the title page and also include collaborating author names as the last paragraph of the “Acknowledgments” section. Please add authors in the format first name, middle initial(s) (optional), and last name. You can add institution or country information for each author if you wish, but this should be consistent across all authors.

8.8 References

Research articles and nonresearch articles (e.g. Review and Commentary articles) must cite appropriate and relevant literature in support of the claims made. Only articles, clinical trial registration records, and abstracts that have been published or are in press, or are available through public e-print/preprint servers, should be cited. References to published books, journal articles, articles in collections, and conference or workshop proceedings, technical reports, and websites should be listed at the end of the manuscript in numbered order of appearance in the text. Any statement in the manuscript that relies on external sources of information (i.e. not the authors’ own new ideas or findings or general knowledge) should use a citation. Authors should give direct references to original research sources whenever possible. Although references to review articles can be an efficient way to guide readers to a body of literature, review articles do not always reflect original work accurately. However, extensive lists of references to original work on a topic can use excessive space. Fewer references to key original papers often serve as well as more exhaustive lists, particularly since electronic literature searching now allows readers to retrieve published literature efficiently. Authors should ensure that their citations are accurate (i.e. they should ensure the citation supports the statement made in their manuscript and authors should not misrepresent another work by citing it if it does not support the point the authors wish to make). Authors should not cite sources that they have not read or preferentially cite their own publications or those of their friends, colleagues, or institution. Authors should also avoid citing work solely from one country. Ideally, authors should cite sources that have undergone peer review wherever possible. Avoid using conference abstracts as references: if unpublished they can be cited in the text, in parentheses. References to papers accepted, but not yet published should be designated as “in press”; authors should obtain written permission to cite such articles and verification that they have been accepted for publication. Information from manuscripts submitted, but not accepted should be cited in the text as “unpublished observations” with written permission from the source. We will generally require copies of manuscripts “in press” or “unpublished observations” to facilitate the review process. Avoid citing a “personal communication” unless it is essential and the information is not available in a public source, in which case the name of the person and date of the communication should be cited in parentheses in the text (See section 8.4.15). Authors must obtain written consent from the person who provided the communication. Excessive and inappropriate self-citation or coordinated efforts among several authors to collectively self-cite is strongly discouraged. Advertisements or adversorial material should not generally be cited unless it is strictly necessary and identified as such.

All references included in the list should be cited in the text using arabic numerals in square brackets before punctuation, so [1], and all references cited in the text should appear in the list. In the reference list, the names of up to the first 6 authors should be cited for each reference, and for articles where there are 8 or more, all other authors are cited as “et al.” meaning “and (more than 1) others”, therefore if there are 7 authors, the 7th author should be cited. The titles of journals should be abbreviated according to the style used for MEDLINE or the U.S. National Library of Medicine (NLM) Catalog of Journals referenced in the National Center for Biotechnology Information (NCBI) Databases (http://www.ncbi.nlm.nih.gov/nlmcatalog/journals). To minimize such citation errors, all references should be verified using either an electronic bibliographic source, such as PubMed, or better PDFs or print copies from original sources and cited exactly
as published (except that titles should be in sentence case). Authors are responsible for the accuracy and completeness of their references and for correct in-text citation. Authors are responsible for checking that none of the references refer to retracted articles except in the context of referring to the retraction. Please do not include DOI or PMID numbers at the end of cited references, unless the article is published online only without bridging page numbers including ahead of print, when DOIs should be given.

8.8.1 In the main text, Tables, and Figure legends

- References should be indicated by numbers in square brackets [thus] in line with the text, numbered consecutively in order of appearance, and placed before punctuation. The actual authors can be referred to, but the reference number(s) must always be given.
- References cited in Tables or Figure legends should be included in sequence at the point where the Table or Figure is first mentioned in the main text.
- Please remove field codes inserted by bibliographic software such as EndNote before submitting manuscripts. This can be done easily by selecting all text (ctrl+a) and using ctrl+shift+f9 on the keyboard.
- Do not cite abstracts unless they are the only available published reference to an important concept. Unpublished abstracts can be cited in the text, in parentheses.
- Do not cite uncompleted work or work that has not yet been accepted for publication (i.e., “unpublished observation”, “personal communication”) as references. See Section 8.4.15.

8.8.2 In the references list

- References should be limited to those cited in the text and listed in numerical order of citation, NOT alphabetical order. Use Vancouver style.
- References should include, in order, authors’ surnames and initials, article title, abbreviated journal name, year, volume (but not month or issue number where pages are continuous throughout a volume) and inclusive page numbers.
- The surnames and initials of all the authors up to 6 should be included, but when authors number 8 or more, list the first 6 authors only followed by “et al.”; where there are 7 authors, include the 7th. Convert given (first) names and middle names to initials for a maximum of 4 initials following each (family) surname.
- Abbreviations for journal names should conform to those used in the U.S. National Library of Medicine, National Institutes of Health, National Center for Biotechnology Information PubMed database.
- If citing a website, provide the author information, article title, website address (uniform resource locator or URL), and the date you accessed the information.
- Reference to an article that is “in press” must state the journal name and, if possible, the year and volume, and DOI.
- Please remove field codes inserted by bibliographic software such as EndNote before submitting manuscripts. This can be done easily by selecting all text and using ctrl+shift+f9 on the keyboard.
- Please remove hyperlinks, vestiges of which are often remaining after cutting and pasting from Internet databases.

Examples of the most common reference types are provided below. (Please pay particular attention to the formatting, word capitalization, spacing, and style including the use of an en dash (–) and not a hyphen (-) to indicate spans of numbers and abbreviated page numbers.) For any types not listed here use the style specified in the Uniform Requirements for Manuscripts Submitted to Biochemical Journals (5th edition) published in the New England Journal of Medicine 1997; 336:309–15 and archived at ICMJE.org or the American Medical Association Manual of Style, 9th edition or later. Another comprehensive source is: Patrias K. Citing medicine: the NLM style guide for authors, editors, and publishers [Internet]. 2nd ed. Wendling D, editor. Bethesda (MD): US National Library of Medicine; 2007- [cited 2018 Jan 5]. Available from: https://www.ncbi.nlm.nih.gov/books/NBK7256/ although we follow this style with variations and options at the discretion of the editors including the retention of diacritics, accent marks, ligated letters and special characters as published, and more than two initials after surnames or family names if published. Hyphenated names as published are indicated with a hyphen or hyphenated initials.

Standard journal articles

Journal supplement
Journal article not in English, but with English abstract (and title)


Journal article not in English (with translated title)


Book with edition


Book with editors


Book chapter in book with editor and edition


Book series with editors


Book chapter in a volume in a series with a separate title and separate authors and editors


Bulletin


Electronic publications


Website


Conference proceedings


Type of article indicated


Theses


Patent


8.9 Tables

Please prepare each Table in MS Word format on a separate page and upload to Editorial manager in order of appearance.
in the text. Tables should be presented without unnecessary vertical or internal horizontal rules (lines). The vertical axis of histograms should not be truncated to exaggerate small differences. Color and shading should not be used. Parts of the Table can be highlighted using superscript, italics, symbols, or bold text, the meaning of which should be explained in the footnote. Please view the American Medical Association Manual of Style (9th edition or later) for further advice on how to prepare Tables for publication.

Tables capture information concisely and display it efficiently; they also provide information at any desired level of detail and precision. An appropriate number of significant figures should be used for all data reported. Including data in Tables rather than text frequently makes it possible to reduce the length of the text. Tables should therefore supplement, not duplicate, the text, and vice versa (the text should not duplicate data presented in Tables). They should have a concise table heading, be self-explanatory, and numbered consecutively in the order of their first citation in the text using arabic numerals and uploaded in that order. Each table should be cited in the text, but the Tables should stand alone and need footnotes to explain any symbols and abbreviations used. Each column should have a short heading title and detailed explanation provided in a footnote if necessary. Items requiring explanatory footnotes should be denoted using superscripted symbols (†, ‡, §, ¶, ††, ‡‡, …), with the footnotes arranged immediately under the Table in alphabetical order. Asterisks (*, **, and hash or pound symbols (#)) should be reserved to indicate the probability level of tests of significance (usually *P < 0.05, **P < 0.01, ***P < 0.001, with #, ##, ### being used for alternative comparisons as is conventional). Abbreviations used in the Table must be defined and placed after the footnotes. If authors include a block of data or table from another source, whether published or unpublished, they must acknowledge the original source and obtain appropriate permission.

8.10 Figures

8.10.1 General guidelines

Figures are critical to display clearly the new knowledge contributed by your work. The number of Figures should be restricted to the minimum necessary to display data to support the text. For clarity and conciseness, it is sometimes helpful to use graphs as an alternative to Tables with many entries. However, do not present the same data in duplicate (e.g., in both graphs and Tables). Each Figure should be cited in the text. Figures should have an informative Figure legend and be numbered in the order of their first citation in the text with an arabic numeral and uploaded in that order. Figure legends should be grouped and placed on a separate page placed at the end of the manuscript following the Reference section. Do not include titles or captions within or on your Figures. The text included in each legend should be sufficient to enable a reader to understand the information in each Figure without reading the main text. All symbols and abbreviations should be defined in the Figure legend. Items requiring explanatory footnotes should follow the same style as that for Tables as described in Section 8.9. Do not use 3-dimensional (3D) graphs where there is no data axis in the third dimension, use a 2D graph instead. The 3D rendering is simply a spurious decorative effect, does not convey meaningful information, and may make interpretation more difficult. Avoid shadows, pixelated computer drawings, and unnecessary aesthetic effects for the same reason. Avoid using shades of gray that differ by less than 30% or colors that are close in hue to identify different symbols or columns in a bar chart.

Digital photographs should be in black-and-white (gray-scale) or in color. Please note that Figures to be printed in black and white should not be submitted in color.

Any information that might identify the hospital or patient, including dates and hospital numbers, should be cropped out, or masked or obscured. All lettering should be made using software and be in proportion to the drawing, graph, or photograph. Photomicrographs must include an internal scale marker, and the legend should state the type of specimen, original magnification, length of the scale marker, and stain.

While it is accepted that authors sometimes need to manipulate images for clarity, manipulation for the purposes of deception or fraud will be seen as misconduct and is not acceptable. The grouping of images from different parts of the same gel, or from different gels, fields, or exposures must be made explicit by using dividing lines (or other graphic means of demarcation) and must also be stated in the Figure legend. No specific feature within an image should be enhanced, obscured (with the exception of for deidentification), moved, removed, or introduced. Adjustments of brightness, contrast, or color balance are acceptable if as long as they do not obscure or eliminate any information present in the original. Nonlinear adjustments (e.g., changes to gamma settings) must be disclosed in the Figure legend.

People with red–green color blindness cannot interpret Figures that use these colors. To quote Okabe and Ito “There is a good chance that the paper you submit may go to color blind reviewers or readers. Supposing that your paper will be reviewed by 3 white males (which is not unlikely considering the current population in science), the probability that at least one of them is colorblind is a whopping 22%! Since
[22% = 1-(1-0.92)^3].” We recommend to provide alternative versions of figures that are more accessible to such individuals and recolor your Figures using magenta instead of red (in the case of double labeling for example with Texas Red and FITC; please see Masataka Okabe and Kei Ito. Color Universal Design (CUD) - How to make figures and presentations that are friendly to colorblind people. [online] 2002 (modified on 2.15.2008, 9.24.2008), Available at http://fly.iam.u-tokyo.ac.jp/color/. Accessed September 03, 2016; or Wong. Nature Methods 2011; 8:441) turquoise and red, or blue and yellow (although this is generally not print friendly). Color schemes that are colorblind safe, print friendly, and photocopier safe can be obtained using tools available at ColorBrewer (v 2.0) found at: colorbrewer2.org

Figures must be submitted as separate picture files at an appropriate resolution (see Section 8.10.2 below). The files should be named according to the figure number and format, and identified with the name of the corresponding author and manuscript reference number (if assigned), e.g., “00035_CorrespondingAuthorName_Fig1.tif”, “00035_CorrespondingAuthorName_Fig2.jpg.” For submission, it is probably helpful for review if multipanel Figures (e.g. with parts A, B, C, D, etc) are submitted as a single correctly-orientated composite that contains all parts of the Figure and uploaded in order of their intended appearance in the text. However, we will accept various panels uploaded separately and this may be required before acceptance. Figures should be closely cropped to minimize the amount of white space around them and facilitate their accurate placement with other elements of manuscripts accepted for publication.

For Asian Biomedicine, the axis labels should be in a sans serif typeface, such as Arial or Helvetica, and will be at least 8 point, preferably 10 pt (2–3 mm), after sizing to one (about 8 cm) or 2 column widths (17 cm) as appropriate. Height should be less than 20 cm. Smaller lettering will yield text that is barely legible. Minimize the number of tick marks on the axes, and do not number each tick if more than 5 ticks. Line weights of about 0.5 pt to 1.5 pt are appropriate. Very thin lines tend to vanish on resizing. Please make sure that all lines are thicker than 0.3 pt when the Figure is sized for publication. All fonts should be embedded. Design Figures so that the least possible number of letters is needed to avoid crowding. Avoid bold lettering, because this looks unpleasantly dark when printed. The style and format of lettering except for indicating subparts of Figures should be uniform throughout all the Figures in a manuscript. Variance of type size within a Figure should be minimal, e.g., do not use 8-pt type on an axis and 20-pt type for the axis label. Use lower case rather than capital letters in the labels of a Figure, but use an initial capital. Leave adequate space between the lettering and the axis. Please also read the American Medical Association Manual of Style (9th or later) for further advice on how to prepare figures for publication. More advice can be found at the following link:

http://art.cadmus.com/da/index.jsp

You may check the suitability of your artwork by using the following free tool http://rapidinspector.cadmus.com/RapidInspector/wi/index.jsp

The Editors reserve the right to redesign or recreate all graphical elements (pictures, graphs, schematic presentations, etc.) of the manuscript if they do not conform to the uniform style of the Journal and may request original data to do so.

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8.10.2 Formats

Regardless of the application used, when your digital artwork is finalized, please “save as” or convert the images to one of the following formats (note the resolution requirements for the final size for line drawings, halftones, and line/halftone combinations given below):

- EPS (Encapsulated PostScript): Vector drawings. Embed the font or save the text as “graphics.” Vector figures should if possible be submitted as PDF files, which are usually more compact than EPS files.
- TIFF (Tagged Image File Format): Color (RGB) or grayscale photographs (halftones) always use a minimum of 300 dpi. TIFF files should be saved with LZW compression, which is lossless (decreases file size without decreasing quality).
- TIFF: Bitmapped line art drawings (purely black and white with no grayscale) use a minimum of 1200 dpi.
- PDF (Adobe Printable Document Format): At appropriate resolution.
- JPEG (Joint Photographic Experts Group): *.jpg at a minimum 600 dpi. JPEG files should be saved at maximum quality.
- CDX (ChemDraw) suitable for molecular structures. Submit as 300 dpi RGB tiff file.
Please do not:

• Supply files that do not meet the resolution requirements detailed above at their final size.
• Supply files that are optimized for screen use (such as GIF, BMP, PICT, WPG) because the resolution is usually too low for print publication.
• Submit graphics that are disproportionately large for their content (please keep file sizes below 10 MB).
• Use Excel to make artwork and avoid the use of MS Office files like PowerPoint or Word.
• Use gratuitous color or red and green combinations in graphs (see point about color blindness above). Color should be used sparingly to identify different categories of data.

8.11 Manuscript checklist (before submission)

□ 1. Title page
  1.1. Corresponding author designated and author(s) to whom correspondence should be addressed after publication of the article if this is different
  1.2. Institutional e-mail address of corresponding author (and author(s) to whom correspondence should be addressed after publication of the article if this is different)
  1.3. Permission to reproduce copyrighted materials
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□ 3. Keywords or short phrases (5–10) (use MeSH where possible)
□ 4. Manuscript text (within specified length limits using IMRAD+C structure)
□ 5. Acknowledgments listed for grants, nonauthorship contributions, technical support, and corporate support. Meeting abstract arising from the work presented. Changes in Address or Affiliation.
□ 6. References double-spaced and cited in the order of appearance
□ 7. Tables (MS Word format)
□ 8. Figure legends
□ 9. Figures (eps, tiff, pdf, jpg)
□ 10. Supporting documents

Of the following items (2), (3), (9), and (10), are mandatory, but may be covered during the submission process. Items (4–8) are required if applicable.

9.4. Copyright permission. See section 13
9.5. Ethics statement. Letter of approval from the relevant ethics review committee. See section 3
9.6. Consolidated Standards of Reporting Trials (CONSORT) flow chart for randomized controlled trials submitted for publication. See section 4
9.7. Statement of documented informed consent. See section 3
9.8. Patient privacy and confidentiality statement. See section 5
9.9. Conflicts of interest statement. See section 2
9.10. Open access agreement

9 The editorial and peer review process

Authors will be provided with a manuscript reference number. If a submission acknowledgment is not received in due course, the author should contact the Editorial Office.

Manuscripts are reviewed by the Editorial Office to ensure that the submission contains all parts and supporting documents. The submission may not be accepted if the author has not supplied all parts of the manuscript and required supporting documents as outlined in these guidelines.

Asian Biomedicine decisions about a manuscript are based on their relevance and importance in an Asian context. All manuscripts must address important scientific questions and the methodology to answer them should be well planned and well implemented, and adhere to internationally accepted ethical and scientific standards (World Association of Medical Editors Publication Ethics Committee. Recommendations on Publication Ethics Policies for Medical Journals, available from: http://www.wame.org/policies). The characteristics of the population and samples (including animal and laboratory samples) must be well defined. Studies with negative results, but sufficient sample size will receive equal consideration. Studies of insufficient sample size are unethical because they put participants at undue risk knowing that the sample size is too small to have sufficient statistical power to answer the questions posed. Outcome measures must be relevant, credible, accurate, and sensitive. Data summaries, statistical analysis procedures, and the interpretations of the results must be appropriate to the study questions.

All studies must be conducted only by scientifically qualified persons and conducted under the supervision of competent clinicians or scientists. Laboratory procedures must be standardized and performed by qualified medical technologists or scientists who have knowledge of the pertinent scientific literature.
Peer review process: The Editor-in-Chief will assess manuscript with the assistance of the Coeditor-in-Chief, Coeditors, and Assistant Editors. If the manuscript is not of sufficient merit to reach priority for publication, does not meet ethical standards, or is not appropriate for the scope of Asian Biomedicine, then it may be rejected without review. At least 2 reviewers, who are appropriate independent experts recruited by a responsible Editor, will review other manuscripts to provide peer review before further processing. The authors will be blinded to reviewers who will be treated anonymously. Where an Editor is on the author list, or has any other competing interest regarding a specific manuscript, another member of the Editorial Board will be assigned to assume responsibility for overseeing peer review. At least one of these reviewers will have no connection with the authors, and will preferably from another country or at least a different institution. If these reviewers disagree on key issues, then further reviewers will be invited to give opinions. The responsible Editor reviews revisions before recommending publication, or not, ahead of final screening. The Editor-in-Chief has the final authority on all editorial decisions.

The Editor-in-Chief and other Editors are experts in various fields. Peer reviewers for manuscripts submitted to Asian Biomedicine are experts in the various topics addressed in the manuscripts chosen by the Editors to provide written assessment of the strengths and weaknesses of the research, with the aim of improving the reporting of the research and identifying the most appropriate and highest quality materials for the journal. They are asked to comment on the relevance of the questions, appropriateness of the methodology, the analysis, the results, and their interpretation. Peer reviewers are asked to state whether the conclusions are consistent with the data provided. They are requested to comment on any ethical concerns raised by studies on animals and humans, and ethical concerns regarding scientific quality. It is hoped they will provide the author with useful and constructive suggestions to improve the manuscript.

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Authors may request exclusion of (oppose) individuals as peer reviewers, but they should explain the reasons in the submission process. Authors should not exclude too many individuals as this may hinder the peer review process. Nevertheless, please note that the Editor may choose to invite excluded peer reviewers.

Intentionally falsifying information, for example, suggesting reviewers with a false name or email address, will result in rejection of the manuscript and is considered serious misconduct. Editors will follow COPE guidelines where this is suspected.

Portability of peer review

To support efficient and thorough peer review, while reducing the burden on peer reviewers, and speeding up the publication process; if a manuscript does not reach the interest criteria of a given journal, but is sound and in scope for Asian Biomedicine, we offer authors the option to transfer past reviewer reports from the other journal with the permission of that journal and previous reviewers.

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The corresponding author will usually be notified within 10 weeks whether the submitted article is accepted for publication, rejected, or subject to further review or revision before acceptance (however, do note that delays are sometimes unavoidable, mostly because of reviewer availability). If revisions are required, authors are asked to return a revised manuscript to the Editorial Office or managing editor within 30 days. Please notify the Editorial Office in advance if additional time is needed. It would be unethical at this stage after review to submit the manuscript for publication elsewhere.

10 Preparation for publication

Once a manuscript has been accepted for publication, authors should ensure the final version of their manuscript in MS Word format has been uploaded, with all Tables and Figures as applicable. The assigned manuscript number should be incorporated into the names of any files.
Accepted manuscripts are then copyedited according to the Journal’s house style and the galley proofs in the form of a PDF file are sent by the managing editor to the corresponding author for final approval. Authors are responsible for all statements made in their work, including changes made by the copy editor.

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