I. EDITORIAL POLICY

The Journal of Nuclear Medicine Technology (JNMT) publishes material of interest to practitioners and scientists in the fields of nuclear medicine and molecular imaging. Proffered articles describing original laboratory or clinical investigations, brief communications, and letters to the editor will be considered for publication. Occasionally, invited articles, editorials, and invited perspectives of selected topics will be published. Manuscripts, including figures and tables, must be original and not under consideration by another publication.

In preparing manuscripts, authors should follow the Recommendations for the Conduct, Reporting, Editing, and Publication of Scholarly Work in Medical Journals (http://www.icmje.org/recommendations/) of the International Committee of Medical Journal Editors and the specific instructions detailed below. Also, helpful guidance in conforming to the Recommendations may be found in Medical Style & Format: An International Manual for Authors, Editors, and Publishers (Philadelphia, PA: Lippincott Williams & Wilkins; 1989) and in the AMA Manual of Style (available by subscription at http://www.amamanualofstyle.com/osocomments.html).

According to the Recommendations, allegation of scientific misconduct or fraud arises if there is substantial doubt about the honesty or integrity of the work, either submitted or published. In the event of allegations of scientific misconduct or fraud, JNMT follows the Recommendations. When appropriate, JNMT reserves the right to present the allegations to the author’s institution or the agency funding the research.

II. MANUSCRIPT SUBMISSION

New manuscripts should be submitted online at https://submit-jnm.snmjournals.org. For instructions visit https://submit-jnm.snmjournals.org/submission/submissionhelp.

Inquiries regarding manuscript status and preparation can be directed to salexand@snmmi.org.

Correspondence about manuscripts should be sent to the JNMT office:
Kathy S. Thomas, MHA, CNMT, PET, FSNMMI-TS
Journal of Nuclear Medicine Technology
Society of Nuclear Medicine and Molecular Imaging
1850 Samuel Morse Drive
Reston, VA 20190-5316
Phone: 703-326-1185
Fax: 703-708-9018

A. Cover Letter

All manuscripts should be accompanied by a cover letter from the author responsible for correspondence about the manuscript. The cover letter should contain a statement that the manuscript has been seen and approved by all authors. If there are more than 10 authors, the specific contribution of each author must be substantiated in the cover letter. The cover letter should inform the editor of potential overlap with other materials already published or submitted for publication and should provide a reference to or a copy of this material. The cover letter should also disclose any conflict of interest—financial or otherwise—that may directly or indirectly influence the content of the manuscript submitted. Finally, the cover letter should provide any additional information that may be helpful to the editor.

B. Authorship, Rights, and Permissions

Each author must have contributed significantly to the submitted work. As recommended by the International Committee of Medical Journal Editors, all authors must have made substantial contributions in all 3 of the following categories:
1. Contributing to conception and design, or acquiring data, or analyzing and interpreting data;
2. Drafting the manuscript, or critically contributing to or revising the manuscript, or enhancing its intellectual content; and
3. Approving the final content of the manuscript.

Simple participation or collection of data alone does not justify authorship but should be mentioned in the acknowledgment section. Changes in authorship after the first review require a written request by the corresponding author and a written authorization from the authors who are to be added or deleted.

If any figures or tables in the manuscript were previously published, this should be acknowledged and written permission from the publisher should be included.

For human studies, approval must be obtained from the institutional review board or equivalent ethics committee and written informed consent must be obtained from research subjects, unless this requirement is waived by the institutional review board or equivalent. For studies in the United States, compliance with the Health Insurance Portability and Accountability Act is also required. Authors must also comply with the clinical trial registration statement from the International Committee of Medical Journal Editors, and the clinical trial registration number must be provided.

For any first-in-human study of a new radiopharmaceutical, the following language should be included in the article to facilitate allowing others to study the drug under the Radioactive Drug Research Committee regulations, rather than having to file additional applications for an investigational new drug or an exploratory investigational new drug: “The mean and standard deviation of the administered mass of [drug] was XX ± YY µg (range, AA–ZZ µg). The mean administered activity was XX ± YY MBq (range, AA–ZZ MBq). There were no adverse or clinically detectable pharmacologic effects in any of the [##] subjects. No significant changes in vital signs or the results of laboratory studies or electrocardiograms were observed [if true].”

For animal studies, approval must be obtained from the appropriate institutional animal care and use committee for compliance with the National Institutes of Health for use of laboratory animals or equivalent.

In compliance with the Copyright Revision Act of 1976, effective January 1, 1978, the following copyright transfer agreement must be faxed, e-mailed, or mailed to the JNMT office. (A printable version is available at https://tech.snmjournals.org/site/misc/ifora.xhtml).

Upon acceptance of the article named above by JNMT, all copyright ownership is transferred to the Society of Nuclear Medicine and Molecular Imaging. We, the undersigned coauthors of this article, have contributed to (1) data design, analysis, or interpretation; (2) writing or critiquing drafts of the manuscript; and (3) approval of the final manuscript before publication. We share in the responsibility for the release of any part or all of the material contained within the manuscript. We also affirm that the manuscript has been seen and approved by all authors. The undersigned warrant that the manuscript (or its essential substance) is new and original, has not been published other than as an abstract in any language or format; and has not been submitted elsewhere for print or electronic publication consideration.

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We also warrant that any human and/or animal studies undertaken as part of the research from which this manuscript was derived are in

INFORMATION FOR AUTHORS 1A
compliance with regulations of our institution(s) and with generally accepted guidelines governing such work.

We further warrant that we have herein disclosed any and all financial or other relationships that could be construed as a conflict of interest and that all sources of financial support for the study have been indicated in the disclosure.

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2. Permission to reprint: The authors retain the following nonexclusive copyrights, to be exercised only after the article has been published in final format in the print version of JNMT.
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   (c) Use the article in theses and/or dissertations.
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   (g) Post a copy of the article on the author’s personal website, departmental website, and/or the university’s intranet, provided a hyperlink to the article on the JNMT website is included.
   (h) In all the instances under clauses 2a through 2g above, the author will give proper credit to the original publication in JNMT as follows: This research was originally published in JNMT. Author(s). Title. J Nucl Med Technol. Year;vol:pp–pp. © by the Society of Nuclear Medicine and Molecular Imaging, Inc.

3. Publish Ahead of Print policy: The authors understand that if and when the manuscript is accepted for publication in JNMT, it will be pre-published online as a Publish Ahead of Print paper. The authors acknowledge that JNMT’s Publish Ahead of Print papers undergo full peer review. Ahead of print articles have not been copyedited, nor have they appeared in a print or online issue of the journal.

Conditions 1–3 in the agreement must be met by all coauthors, and the agreement must be signed by all coauthors. Designate “first author” and “corresponding author” in parentheses by their signatures.

This copyright transfer requirement does not apply to work prepared by U.S. government employees as part of their official duties. However, the authors of such work are required to complete section B of the copyright transfer agreement found at https://tech.snmjournals.org/site/misc/cfora.xhtml.

III. MANUSCRIPT REVIEW, REVISION, AND RESUBMISSION

Submitted manuscripts are reviewed for originality, significance, adequacy of documentation, composition, and adherence to these guidelines. However, editorial decisions are based not only on the technical merits of the work but also on factors such as priority for publication and relevance to the general readership of JNMT. All manuscripts are judged in relation to other submissions currently under consideration.

Two helpful publications to read before writing a manuscript are “The Efficacy of Diagnostic Imaging” by Fryback and Thornbury (Med Decis Making, 1991;11:88–94) and “Bias in Research Studies” by Sica (Radiology, 2006;238:780–789). At the discretion of the Editors, the manuscript may be returned rapidly—without external peer review—if deemed not competitive or outside the scope of JNMT. Rebuttals to rejected manuscripts are strongly discouraged, and requests for resubmission of rejected manuscripts are generally not granted without significant demonstration of errors in the review or decision process. Most articles are rejected on grounds of insufficient priority or lack of relevance to JNMT, not data quality or technical issues.

Manuscripts must be written in English. When necessary, authors should seek the assistance of experienced, English-speaking medical editors before submission. A medical editor should review the final draft of the original and any revisions of the manuscript. Authors will be required to provide revisions of articles written in substandard English before peer review.

Manuscripts considered suitable for review are evaluated by 2 reviewers. The Editors select the reviewers and make the final decision on the manuscript. Authors may suggest reviewers for their manuscripts. Referees who review a manuscript remain unknown to the authors.

It is unusual for a manuscript to be accepted for publication without first undergoing a process of revision. Revised manuscripts are judged on the adequacy of responses to suggestions and criticisms made during the initial review. Revision of a manuscript does not guarantee acceptance. A revision should be accompanied by a point-by-point reply to the reviewers’ and editors’ critiques in which any changes are briefly described. The authors also should provide justification for not altering the manuscript in response to any reviewer comments believed to be inappropriate. Red font should be used to indicate all changes within the manuscript itself, and a clean version of the manuscript should be provided.

The revised manuscript and accompanying reply must be submitted to JNMT via the online submission and review website at https://submit-jnm.snmjournals.org within 30 days of the date of the editorial decision. If circumstances prevent completing the revisions by the deadline, please contact Susan Alexander at 703-326-1185 or at salexand@snmmi.org. If the revisions are not received within 3 months after being requested, the manuscript may be started on a new review cycle and given a new manuscript number.

All accepted manuscripts are subject to editing for accuracy, clarity, and style.

IV. ARTICLE TYPES

Original scientific and methodology articles should contain no more than 6,000 words. This word limit includes all data: title page, abstract, text, disclosure, acknowledgments, key points, references, figure legends, and tables. The goal is to limit original articles to 8 printed pages. A maximum of 7 figures (maximum of 14 parts in total with no more than 4 parts per figure preferred), 7 tables, and 40 references is allowed. Abstracts should be structured (see VI. C. below) and should contain a maximum of 350 words.

Teaching case studies should contain no more than 1,000 words. This word limit includes all data: title page, abstract, text, disclosure, acknowledgments, references, figure legends, and tables. A maximum of 5 figures and 10 references is allowed, and the maximum number of authors is three. The objective of teaching case studies is to present images that demonstrate key facts or concepts in clinical nuclear medicine and molecular imaging. Emphasis is placed on studies in which imaging has been useful in helping with the diagnosis. Teaching case studies will be accepted for publication in JNMT at the discretion of the Editor and may also be posted on the SNMMI website. Submissions not accepted for publication in JNMT may be accepted for posting on the SNMMI website only. Teaching case studies should include a brief, unstructured abstract followed by 4 sections: an “Introduction” section briefly describing the point that the article is teaching, explaining the significance of the article, and summarizing its educational value; a “Case Report” section describing relevant medical history, laboratory findings, clinical course, procedures
performed, and condition at last follow-up; a “Discussion” section describing any findings, differential diagnosis, and final diagnosis; and a “Conclusion” section summarizing the take-home teaching points.

**Quality case studies** should contain no more than 3,000 words. This word limit includes all data: title page, abstract, text, disclosure, acknowledgments, figure legends, and tables. A maximum of 4 figures, 2 tables and 20 references is allowed, and the maximum number of authors is three. The objective of a quality case study is to present a specific event that required an in-depth analysis resulting in an operational assessment, resolution, corrective action, and verification of effectiveness using standard quality assessment tools (e.g., a root cause analysis using Lean or Six Sigma methodology). The quality case study should contain a brief, unstructured abstract followed by 4 sections: an introduction (a description of the event, answering the questions what, how, when, where, and who), a “Quality Analysis” section (repeatedly asking the question why until a solution to the error is found), a “Corrective Action” section, and a “Verification of Effectiveness” section.

**Letters** should concern previously published material or matters of general interest and should be brief and to the point. Letters should be given a title and should also be accompanied by a copyright transfer agreement as specified above in Manuscript Submission. All material is subject to editing. Letters commenting on previously published articles should be received within 1 year of the date of the referenced article’s publication. Letters should contain no more than 800 words; this word limit includes all data: title page, authors and affiliations, the letter itself, and any references. Letters should contain no figures or tables and no more than 10 references.

**C. Abstract**

A structured abstract must be included with each original scientific manuscript, including brief communications. The abstract should contain a maximum of 350 words for original scientific and methodology articles or 150 words for brief communications and include 4 clearly identifiable elements of content: rationale (goals of the investigation), methods (description of study subjects, experiments, and observational and analytic techniques), results (major findings), and principal conclusions. Except for the rationale, these sections should be preceded by headings (i.e., **Methods, Results, and Conclusion**). Three to 5 key words should also be submitted with the abstract.

A graphical abstract must also be included with each original scientific manuscript, including brief communications. The graphical abstract, a visual aid to understanding your key findings, is intended to attract the attention of readers and help them find articles that interest them. It should be clear, concise, eye-catching, and easily understood by a typical *JNMT* reader. It will be displayed in the online article and online table of contents and might also be used in promotional media, in the *JNMT* carousel, or on the *JNMT* cover if your article is selected to be featured in that issue. It must comprise a single, simple, original graphic that is not densely packed with information, has no legend, and is not a figure from your article. It cannot be a table, be reprinted from another source, or include any trade names, logos, or images of trademarked items. Text can be used sparingly but must be in Arial or Roboto font large enough to be legible. For ease of comprehension, it should have a clear start and end, with the information preferably running from top to bottom or left to right. Build the graphical abstract using the PowerPoint
INFORMATION FOR AUTHORS

D. Text

Describe procedures in sufficient detail to allow other investigators to reproduce the results. Do not use hyperbolic terms or phrases in the title, abstract, or body of the text. Qualitative claims as to the superiority (superior, best) or primacy (first, novel, unique) of an idea or instrument are not acceptable. Do not use numbered or bulleted lists. Any brand-name or trademarked instrument, pharmaceutical, or other product mentioned must be followed by the name of the manufacturer, in parentheses. The use of generic names is preferred to the use of brand names or trademarked names. Original scientific and methodology articles are divided into the following sections:

Introduction
This section should be brief and focused. The final paragraph should state the hypothesis investigated.

Materials and Methods
This section should include statements about Institutional Review Board approval, written informed consent, compliance with the Health Insurance Portability and Accountability Act, and animal care committee approval, as appropriate. The standard statement for institutional board approval and consent is the following: “The study has been approved by the institutional review board [or equivalent], and all subjects signed an informed consent form [or the need for written informed consent was waived].” The clinical trial registration number should also be included when appropriate (http://www.clinicaltrials.gov/). For any first-in-human study of a new radiopharmaceutical, the mean, standard deviation, and range of the administered mass of drug and mean administered activity need to be provided, as well as clinically detectable pharmacologic effects. The checklist and flow diagram from one of the following evidence-based statements should be followed as appropriate and submitted as supplemental material: STARD (http://www.stard-statement.org); CONSORT (http://www.consort-statement.org); REMARK (http://www.nature.com/nrclinonc/journal/v2/n8/full/nponc0252.html). The number and selection of subjects must be clearly described, as well as the prospective or retrospective nature of the study. Procedures must be described in enough detail to allow reproducibility by others. The last paragraph should describe the statistical methods.

Results
The text of this section should not repeat information presented in the tables and figures. When percentages are given, the ratio of numerator to denominator should be in parentheses.

Discussion
This section should summarize any advances in knowledge provided by the results and then discuss their implications in relation to other studies. Limitations and biases of the study must be addressed. The direction of future research may be mentioned.

Conclusion
This section should be brief, should summarize the key points of the paper, and should not introduce new material or references.

E. References

References (not to exceed 40 in an original scientific or methodology article, 20 in a brief communication, 10 in a letter, or 5 in a teaching case study) should be cited in consecutive numeric order at first mention in the text and designated by reference number italicized, in red font, and in parentheses. References appearing only in a table or figure should be placed at the end of the reference list.

When listing references, follow the AMA Manual of Style: A Guide for Authors and Editors (available by subscription at http://www.amamanualofstyle.com/osopublic/index.html). Abbreviate journal names according to the journals database available at PubMed.gov. For journal articles, include the year and volume number in the citation but not the month or issue number. “Unpublished observations” and “personal communications” should not be used as references, although written or oral personal communications may be noted as such in the text. References cited as “in press” must have been accepted for publication and not merely be in preparation or submitted. The author is responsible for the accuracy of all references and must verify them.

List all authors when 6 or fewer; for 7 or more, list the first 3 followed by “et al.”

Examples of journal articles:

Example of a book and book chapter:

Example of an Internet reference:

Example of a package insert:

Authors are encouraged to generate their references using EndNote (Thomson Scientific). The JNMT Output Style for EndNote is available at http://endnote.com/downloads/styles.

F. Units of Measurement

All measurements should be listed in Système International (SI) units. Non-SI units may be used after the SI units but should be placed in parentheses. Use becquerels, not curies, as the unit of activity (1 mCi = 37 MBq).

G. Abbreviations and Symbols

With the exception of units of measurement, JNMT strongly discourages the use of abbreviations. Whenever possible, terms should be spelled out in full rather than being abbreviated. Every abbreviation, even those that are well known and in common use, must be defined the first time it is mentioned in the manuscript; spell out the full term and place the abbreviation, in parentheses, after the full term.

H. Tables

Place tables at the end of the manuscript file; do not submit them as separate files. Do not submit tables as images. Tabbed or space-separated table text is not allowed; tables should be created in Microsoft Word table format or a similar format. The
The number of tables is limited to 7, except in the case of dosimetry articles, which may exceed that number in lieu of number of tables. Use vertical rules. Give each column a brief heading. The title and column headings and at the end of the table. Do not duplicate, the text. Each table must be cited in consecutive numeric order in the text. Number the tables consecutively with an Arabic numeral after the word "TABLE." Titles should be descriptive and brief. Horizontal rules should be placed below the title and column headings and at the end of the table. Do not use vertical rules. Give each column a brief heading.

Place explanatory matter in footnotes, not in the title. Use the following symbols in this sequence: *, †, ‡, §, ¶, ‖, ‖, ‖, **. In a footnote, define all abbreviations in the order in which they appear in the table and identify statistical measures of variations, such as standard deviation and standard error of the mean. If data from another published source are used, obtain written permission from the publisher, cite the original source in the references, and include the following credit line in a footnote: “Reprinted with permission of Ref X.” If data from an unpublished source are used, obtain permission from the principal investigator and acknowledge fully.

**AQARA Requirements for Radionuclide-Based Images**

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*%IA/cm³ = percentage of injected activity per cubic centimeter of tissue.*
or .pdf. If the .jpeg file format is used, the images must be of medium quality or better (quality setting of at least 6). Each figure, including those in .ppt, .doc, and .pdf formats, must be submitted as a separate file. Each figure must also be included in the manuscript file before its respective legend. Crop and size digital figures to match figure specifications and to minimize total file size.

All submitted illustrations become the property of the Society of Nuclear Medicine and Molecular Imaging.

Videos can be published as supplemental data online.

J. Figure Legends

Legends for figures should be concise and should not repeat the text. Number the legends with an Arabic numeral after the word “FIGURE.” If a figure has more than one part, each part clearly. Any letter designations or arrows appearing on the figures should be identified and described fully. Abbreviations used in each figure should be defined in the legend in alphabetical order.

Besides being submitted as a separate file, each figure should be inserted before its respective legend in the figure legends section of the manuscript file.

Original (not previously published) figures are preferred for publication in JNMT; however, if figures have been published previously, authors are responsible for obtaining written permission from the publisher to reprint. The source of the original material must be cited in the references and the following credit line in parentheses included in the legend: “Reprinted with permission of Ref. X.” All permission letters should be submitted online at the time of manuscript submission.

K. Disclosure

A statement of disclosure is required for all submissions. Include in the disclosure any potential conflicts of interest as reported in the disclosure form of the International Committee of Medical Journal Editors. For the work under consideration for publication, these include any payments received from a third party, such as grants, consulting fees, travel fees, or honoraria. Also disclose any relevant financial activities outside the submitted work, such as employment, royalties, stock options, or patents. For industry-sponsored or industry-supported research, authors who are not employees of or consultants to the industrial entity must be specified as having control of any data that might present a conflict of interest for employees or consultants. If no conflicts exist, this should be specified in the statement.

L. Acknowledgments

Individuals whom the authors wish to thank may be listed in the acknowledgments. In addition, persons who have contributed intellectually to the work but do not fulfill conditions 1–3 of the copyright transfer agreement in section II may be listed.

M. Key Points

Original submissions must include 3 key points just before the reference list:

KEY POINTS

QUESTION: A one-sentence focused question based on the study hypothesis or goal.

PERTINENT FINDINGS: One or two sentences on the design (e.g., clinical trial, cohort study, case-control study, meta-analysis), the primary outcome, and the findings (only basic numbers and whether they are statistically significant, but not the results of statistical tests or measures of variance).

IMPLICATIONS FOR PATIENT CARE: One sentence on how the findings might affect patient outcomes or, for basic science, what the translational implications might be.

N. Supplemental Data

All data that are needed to support the central conclusions of the article must be presented in the manuscript itself. Other data (e.g., large-scale tabulations) that are integral to the manuscript and of interest to specialists but not practical to include in the printed journal can be submitted for online-only publication as supplemental data. The data may include images with legends, tables, or videos; supplemental text is discouraged (if some of the methods have been described in a previous publication, the manuscript can reference that publication). Each item of supplemental data should be given a brief descriptive title and should be directly referred to in the manuscript (e.g., Supplemental Table 1). Because supplemental data files are placed online unedited, as submitted by the author, the uploaded files need to be final and ready for publication. Provide original files rather than .pdfs. Do not include a title page.

VI. CHECKLIST FOR NEW SUBMISSIONS

___ Is all text in the manuscript double-spaced, including the references?
___ Does the title page include the title, short running title, and authors’ names and complete affiliations; complete address, telephone number, fax number, and e-mail address for the first author, if different from the corresponding author?
___ Are the figures and tables in consecutive numeric order?
___ Have the figures been included in the manuscript file before their respective legends, as well as being submitted as separate image files of an acceptable format and resolution?
___ Do all PET, SPECT and planar nuclear images follow AQARA requirements (http://jnm.snmjournals.org/site/misc/AQARA.xhtml)?
___ Has permission been obtained from the publisher to reprint previously published figures and tables?
___ Has the copyright transfer agreement been signed by all authors?
___ Was the study approved by an institutional review board or equivalent, and has this approval been mentioned in the “Materials and Methods” section?
___ Did all subjects give written informed consent, or did the institutional review board waive the need to obtain informed consent?
___ Was the study approved by an institutional animal care and use committee or equivalent?
___ Has the clinical trial registration number been provided?
___ Has first-in-human radiopharmaceutical language been included?
___ Did you follow the checklist and flow diagram from one of the following evidence-based statements, and did you submit the checklist as supplemental material: STARD (http://www.stard-statement.org); CONSORT (http://www.consort-statement.org); PRISMA (http://www.prisma-statement.org/statement.htm); REMARK (http://www.nature.com/nrclinonc/journal/v2/n8/full/ncponc0252.html).