

CALL FOR ABSTRACTS

May 30-June 3, 2025 Chicago, IL & Online am.asco.org

ABSTRACT SUBMISSION DEADLINE:

JANUARY 28, 2025, AT 11:59 PM (ET)

The 2025 ASCO Annual Meeting is The Place to Be Heard

We are excited to bring the global oncology community back together in Chicago to foster new connections and discover the latest innovative findings in the study, diagnosis, and treatment of people with cancer. Plan to participate in the world's largest clinical cancer research meeting—prepare your abstract now and submit it by January 28.

Key Dates

NOVEMBER 6, 2024	Abstract Submission Opens
EARLY DECEMBER 2024	ASCO Member Registration and Hotel Reservations Open
MID-DECEMBER 2024	Registration and Hotel Reservations Open for All
JANUARY 28, 2025, AT 11:59 PM (ET)	Abstract Submission Deadline
MARCH 5, 2025, AT 12:00 PM (ET)	Late-Breaking Abstract Submission Deadline (Shell Submission Required by January 28)
MARCH 25, 2025	Abstract Notifications Sent to First Authors
APRIL 6, 2025	Abstract Withdrawal Deadline
APRIL 23, 2025, AT 10:00 AM (ET)	Regular and Late-Breaking Abstract Titles Released Online
APRIL 23, 2025, AT 11:59 PM (ET)	Hotel Reservation and Early Registration Deadline
MAY 22, 2025, AT 5:00 PM (ET)	Regular Abstracts Released Online
MAY 30-JUNE 3, 2025	2025 ASCO Annual Meeting at McCormick Place Late-Breaking Abstracts (LBAs) release on day of presentation at the meeting

Highlights from the 2024 ASCO Annual Meeting



45,000 Attendees, with 87% attending in person



920M X/Twitter impressions of #ASCO24 hashtag from May 31-June 4, 2024



Garnered more than 17,000 news articles reaching an audience of +1 billion



"Knowledge is a powerful force in advancing progress against cancer when intentionally applied. The ASCO Annual Meeting is the premier global forum for exchanging the latest high-impact clinical cancer research and learning how to implement these advances at the bedside. By sharing your scientific

discoveries at our Meeting, you will equip our patients and the oncology community with timely, evidence-based research while inspiring equitable care delivery around the world."



"ASCO's Annual Meeting promises something for everyone who is passionate about advancing care and treatment for people facing cancer from fantastic science and clinical trial presentations, to educational sessions updating us on current and emerging therapies."

> - Erika Hamilton, MD Chair, 2025 ASCO Scientific **Program Committee**

Submit Your Research to #ASCO25

As you prepare your submission to the Meeting, please make note of the following details:

- All types of oncology-related research are eligible for submission. Please note case reports are not accepted.
- · Abstract should address scientific questions, detail clinical observations, or contain primary scientific data.
- Data from the long-term follow-up of previously presented clinical trials may be submitted only if significant new information can be shown. Authors will be required to provide details about what new data is included in the ASCO submission.
- In general, ASCO discourages reporting of interim results unless approved by the study's Data and Safety Monitoring Committee. Please review ASCO's Abstract Biostatistical Guidelines for additional information.

- · Abstracts of clinically related subjects should be combined into a single abstract. Submission of multiple abstracts on a single study may result in rejection of one or more abstracts.
- To ensure the integrity of the review process, corrections to abstracts will not be allowed after the abstract submission deadline.

Simultaneous Publication in **ASCO Journals**

Authors of high-impact, practice-changing studies who are interested in having their research published simultaneously at the meeting should submit a pre-submission inquiry to jco@asco.org.

Getting Started: To submit an abstract, you will need to log in with an ASCO.org account. If you are not an ASCO member, you can create a guest account. The person submitting the abstract is not required to be an author on the abstract and will be able to select the first author on the designated step. However, the first author will need to agree to all ASCO policies and will be held responsible for any violation of the policies.



Don't Miss the Opportunity to Present your Research at #ASCO25

- **Plenary Session**
- **24** Oral Abstract Sessions
- Clinical Science Symposia, including 3 Special **Cross-Cutting Symposia**
- Rapid Oral Abstract Sessions
- Poster Sessions, including Trials in Progress

Over 3.100 Total Presentation Opportunities



The Place to Be Published

Author and Sponsor Eligibility

Authorship

As you prepare your submission to the Meeting, please make note of the following eligibility criteria:

- · Individuals may submit up to two regular abstracts as the first author.
- Individuals may submit an unlimited number of Trials in Progress abstracts.

Sponsorship

ASCO membership is not required to submit an abstract; however, each abstract must be sponsored by an ASCO member*.

- New in 2025: All ASCO members in good standing may sponsor their own submissions.
- Abstract submitters who are not ASCO members must have their abstract sponsored by an ASCO member. There is no limit to the number of abstracts a member can sponsor.
- Abstract sponsors must verify the contents of the abstract and support its data and comply with all Abstract Policies.

*Review ASCO's Member Benefits or contact ASCO Customer Service at 703-299-0158 or 1-888-282-2552 for more information.

Submission Tracks and Subcategories

The 2025 ASCO Annual Meeting Scientific Program Committee seeks abstracts in the following categories. Authors will be asked to select a track and subcategory when submitting their abstract.

Breast Cancer-Local/Regional/Adjuvant

Adjuvant Therapy Biologic Correlates DCIS/LCIS/Premalignant Lesions **Local-Regional Therapy Neoadjuvant Therapy**

Breast Cancer-Metastatic

Biologic Correlates HER2-Positive Hormone Receptor-Positive Triple-Negative Non-Subtype-Specific Breast **Cancer Therapies Other Breast Cancer Subtypes**

Care Delivery/Models of Care

Access to Care Care Delivery Clinical Informatics/Advanced Algorithms/Machine Learning **Digital Technology Disparities in Care** Geriatric Models of Care **Global Models of Care Other Models of Care Survivorship Models of Care Telemedicine/Remote Care**

Central Nervous System Tumors

Brain Metastases Primary CNS Tumors-Glioma Primary CNS Tumors-Non-Glioma

Developmental Therapeutics-Immunotherapy

Antibodies Cellular Immunotherapy Circulating Biomarkers Immunobiology New Targets and New Technologies (IO) PD1/PD-L1 Inhibitor Monotherapy PD1/PD-L1 Inhibitor Combinations

Tissue-Based Biomarkers Vaccines

Other Checkpoint Inhibitors (Non- PD1/ PD-L1, Monotherapy, or Combination) **Other IO-related Topics**

Developmental Therapeutics-Molecularly Targeted Agents and Tumor Biology

Chemotherapy and Antibody-**Drug Conjugates Circulating Biomarkers Molecular Diagnostics and Imaging New Targets and New Technologies** (Non-IO)

Pharmacology/Pharmacodynamics/ **Pharmacogenetics**

Radiopharmaceuticals **Small Molecules Tissue-Based Biomarkers** Other Developmental Therapeutics

Gastrointestinal Cancer-**Colorectal and Anal**

Anal Cancer Epidemiology/Outcomes Colorectal Cancer-Advanced Disease Colorectal Cancer-Local-Regional Other Colorectal and Anal Cancer

Gastrointestinal Cancer-Gastroesophageal, Pancreatic, and Hepatobiliary

Esophageal or Gastric Cancer-Advanced/Metastatic Disease Esophageal or Gastric Cancer-

Local-Regional Disease Hepatobiliary Cancer-

Advanced/Metastatic Disease

Hepatobiliary Cancer-Local-Regional Disease Neuroendocrine/Carcinoid

Pancreatic Cancer-Advanced/Metastatic Disease

Pancreatic Cancer-**Local-Regional Disease Other GI Cancer**

Genitourinary Cancer— Kidney and Bladder

Kidney Cancer Urothelial Cancer-Advanced/Metastatic Urothelial Cancer-Localized Disease Other Kidney and Bladder Cancer

Genitourinary Cancer-Prostate, Testicular, and Penile

Epidemiology Outcomes Germ Cell/Testicular **Penile Cancer**

Prostate Cancer-Advanced/ **Castrate-Resistant**

Prostate Cancer-Advanced/ **Hormone-Sensitive**

Prostate Cancer-Local-Regional Disease Other Prostate, Testicular, and **Penile Cancer**

Gynecologic Cancer

Cervical Cancer Ovarian Cancer Uterine Cancer Other Gynecologic Cancer

Head and Neck Cancer

Advanced Disease Biologic Correlates Local-Regional Other Head and Neck Cancer (Salivary, Thyroid)

Hematologic Malignancies-Leukemia, Myelodysplastic Syndromes, and Allotransplant

Acute Leukemia **Allogenic Stem Cell Transplantation** Chronic Leukemia—CML Myelodysplastic Syndromes (MDS) Myeloproliferative Syndromes (MPD) and Mast Cell Disorders Other Leukemia, Myelodysplastic

Syndromes, and Allotransplant

Submission Tracks and Subcategories, continued

Hematologic Malignancies-**Lymphoma and Chronic** Lymphocytic Leukemia

Cell Therapy, Bispecific Antibodies, and **Autologous Stem Cell Transplantation** for NHL, HL, or CLL

Chronic Lymphocytic Leukemia (CLL) and Hairy Cell

Hodgkin Lymphoma Non-Hodgkin Lymphoma **Other Lymphoma**

Hematologic Malignancies-Plasma **Cell Dyscrasia**

Cell Therapy, Bispecific Antibodies, and **Autologous Stem Cell Transplantation** for Plasma Cell Disorders

Multiple Myeloma **Plasma Cell Disorders**

Lung Cancer-Non-Small Cell Local-Regional/Small Cell/Other **Thoracic Cancers**

Adjuvant Therapy Biologic Correlates Local-Regional Non-Small Cell Lung Cancer Mesothelioma **Small Cell Lung Cancer Thymic Malignancies**

Lung Cancer-Non-Small Cell Metastatic

Biologic Correlates

Targeted (Non-Immunotherapy) **Immunotherapies**

Medical Education and Professional Development

Clinician Burnout and Wellness Education Research Professionalism and Ethics Social Media Research Workplace Disparities/Issues

Melanoma/Skin Cancers

Advanced Disease Local-Regional Other Melanoma/Skin Cancers

Pediatric Oncology

Leukemia/Lymphoma **Pediatric Solid Tumors** Survivorship

Symptom Management/Supportive Care/ **Palliative Care**

Prevention, Risk Reduction, and Genetics

Cancer Prevention Etiology/Epidemiology **Germline Genetic Testing Health Promotion/Behaviors Hereditary Cancer Syndromes**

Quality Care/Health Services Research

Clinical Research Design Health and Regulatory Policy Health Outcomes Health Services Research Patient Reported Outcomes Quality Improvement Real-World Data/Outcomes

Sarcoma

Bone Tumors Gastrointestinal Stromal Tumors (GIST) Molecular Targets/Biomarkers/ **Tumor Biology Soft Tissue Tumors**

Symptom Science and Palliative Care

Cardio-Oncology **End-of-Life Care** Late and Long-Term Adverse Effects **Palliative Care and Symptom Management Pathobiology of Symptoms** Psychosocial and **Communication Research** Toxicities-Prevention and Management **Strategies**



am.asco.org/submission-tracks-categories

Late-Breaking Abstract (LBA) Submission Guidelines

The ASCO late-breaking abstract policy allows for the submission of late-breaking data only for:

 Randomized phase II and III trials for which no preliminary data are available at the time of the abstract submission deadline (January 28, 2025);

OR

 Original research studies that highlight novel and high-impact research with practice-changing implications.

The initial late-breaking placeholder (shell) must be submitted by the January 28 deadline. Final late-breaking data will be due by March 5, 2025.

During submission, you will be required to provide:

- · the primary clinical endpoint for analysis
- · type of analysis
- · date of planned analysis
- · planned statistical methods for analysis

The late-breaking data policy is not a mechanism to allow for updated data to be submitted later when preliminary data are available by the abstract submission deadline.

Phase III clinical research trials for which the final data are not available by the March-5 deadline may be granted an extension to submit; however, the initial trial information MUST be submitted by the January 28 deadline. Contact abstracts@asco.org with questions.

am.asco.org/late-breaking-data-submission-guidelines



Trials in Progress (TPS) Abstract Submission Guidelines

ASCO recognizes the importance of bringing together researchers to discuss ongoing trials. Trials in Progress posters provide an opportunity for members of the research community to present ongoing trials, foster collaboration, and discuss correlatives and novel trial designs. In addition, Trials in Progress highlight the transition of emerging biologic pathways and new agents into the clinic-providing "coming attractions" for oncologists in clinical practice.

All phases of clinical research (phases I to III, supportive care, nonpharmacologic interventions) may be considered for inclusion as a Trials in Progress submission. Trials submitted to this session are ongoing and have not reached pre-specified endpoints

for analysis. As such, inclusion of results would be improper and is strictly forbidden.

To maximize visibility and communication around ongoing trials, Trials in Progress abstracts are excluded from ASCO's Prior Presentation/Publication and Confidentiality Policies but must adhere to and abide by the Copyright Transfer Policy. If copyright has been assigned to another entity, the first author must obtain permission from the copyright holder for ASCO to reprint the abstract. Authors are required to notify abstracts@asco.org and provide appropriate copyright language from the other entity, should their Trials in Progress abstract be published before ASCO publishes the abstract.

Abstracts should be organized according to two sections, Backgrounds and Methods, as described below:

Background

- · Scientific background/rationale for the trial.
- · Preclinical and/or earlier-phase clinical data that have already been publicly presented or published may be included with references. The Trials in Progress abstract should not be used to present preclinical or earlier-phase clinical data for the first time.
- Correlative studies of particular interest.

Methods

- · Trial design and statistical methods, highlighting any novel aspects of the design.
- · Treatment or intervention planned.
- Major eligibility criteria, highlighting unusual aspects.
- · Current enrollment without providing results or endpoints.
- · Clinical trial registry number (required).
- · Examples:
 - Phase I studies may say, "Cohorts 1 and 2 have been completed without DLT. Enrollment to cohort 3 began in January 2025".
 - Phase II studies may report, "8 of planned 32 patients have been enrolled" or "Prespecified activity goal for the first stage of accrual was met; second stage accrual began in January 2025".
 - · Phase III trials may report, "The DMC last reviewed the trial in December 2024 and suggested that the trial continue as planned".
 - Enrollment must have already begun or have been completed with no data analysis available by the submission deadline (there are no exceptions to this criterion).

The following information is not acceptable in a Trials in Progress abstract and/or poster:

- · Any preliminary data including toxicity, response rate, pharmacokinetic, or correlative analyses. Abstracts including results or preliminary data will be rejected without further review.
- · Proprietary drug names or the names of drug manufacturers in the title or body of the abstract. If necessary, you may include the proprietary drug name in parentheses directly after the generic name on first use in the body of the abstract. ASCO reserves the right to replace proprietary names with generic names to adhere to this requirement.
- · Information about pricing, fees, or reimbursement related to trial participation.



Abstract Submission Requirements

As you prepare your abstract submission to the Meeting, please make note of the following requirements.

- Identification of Original Research: Indicate whether your abstract reports on original research. Original research means a systematic investigation designed for the purpose of expanding knowledge or understanding, including the analysis of data. For clarity, a clinical trial is original research under this definition, and a summary or review of prior knowledge is not original research under this definition.
- Identification of Clinical Trials: Indicate whether your research is a clinical trial. A clinical trial is a research study in which one or more human subjects are prospectively assigned to one or more interventions (which may include placebo or other control) to evaluate the effects of those interventions on healthrelated biomedical or behavioral outcomes (National Institutes of Health [NIH] Office of Extramural Research). Though clinical trial registration is not required for abstract submission, publication, or presentation, certain clinical trials are required to be registered by law and/or prior to journal publication. If a clinical trial is already registered, the first author will be asked to provide the name of the registry and the trial registration number during the abstract submission process. The clinical trial number will be included as part of the published abstract.
- Funding Source: Indicate the abstract funding source(s). The funding sources entered here will be published with your final abstract. They do not need to be included within the abstract body.
- Abstract Title: The title should objectively describe the study and be written in sentence case. Titles should not allude to a result or conclusion of the study. ASCO reserves the right to edit conclusive titles.
- Track and Subcategory: Select the most appropriate track and subcategory for the abstract. When submitting

- your abstract, you will have the option of identifying a secondary subcategory for your abstract. The ASCO Scientific Program Committee has the authority to recategorize an abstract.
- Coauthor(s): Provide the full name, academic degree(s), institution, address, email address, and disclosure information for each author. You may list up to 20 individual authors for each abstract. Ensure that the correct author information and order is submitted. Changes to author names or order will not be accepted after the submission deadline.
- Abstract Body and Table: The body of the abstract should describe the background, methods, results, and conclusions of the research. Type the abstract directly into the text box or cut and paste from an existing document.
 - · Character count: Do not exceed 2,600 characters including the abstract title, body, and table. The character count does not include spaces, author names, or institutions.
 - Table: One data table is permitted per abstract. The composition process does not enable shading or the merging of cells with centered text. Limit the table to no more than 10 rows. Illustrations and figures are not permitted.
 - ASCO discourages the submission of complex tables with subrows and subcolumns. Tables that do not render correctly in print or online will not be fixed after publication.
- Submission Fee: An \$80 (USD) nonrefundable submission fee will be charged per abstract submitted. Payment is due at the time of submission. Credit cards are the only accepted form of payment. Checks, wire transfers, and purchase orders will not be accepted.
 - Payment waivers: ASCO is pleased to offer payment waivers to first authors from low-income countries and lower-middleincome countries (LMICs), as defined by the World Bank. Authors may apply for a payment waiver at the time of submission. Please contact us for more details.
 - · Please note: The abstract submission fee does not include registration for the Annual Meeting.

am.asco.org/abstract-submission-requirements

Abstract Submission Requirements, continued

Save Time, Ensure Accuracy—Collect Author Disclosure Information Early

- Disclosure Declaration: ASCO's policy promotes balance, independence, objectivity, and scientific rigor in all its activities through the disclosure of financial interests and other relationships, and management of potential conflicts. The financial interests or relationships requiring disclosure are outlined in ASCO's Policy For Relationships With Companies (Journal of Clinical Oncology 2017 35:7, 796-798). All authors are expected to disclose all relationships with for-profit health care companies.
 - · The Coauthor Disclosure Form may be used by the first author to obtain disclosure information from coauthors. The submitter must enter all disclosure information through the Abstract Submitter by the submission deadline.
 - · If an author has provided disclosure through the ASCO Disclosure Management System (coi.asco.org), the information will automatically populate in the submission site.

- · Presenter Eligibility: At least one coauthor on the abstract must be eligible to present. If the first author is employed by a company or holds ownership with a private company (see company definition below), an alternate presenter who does not have a relevant employment or ownership relationship must be named from the list of coauthors. This applies to abstracts presented in an oral abstract session, rapid oral abstract session, or clinical science symposium.
 - · Company is defined as an entity whose business is developing, producing, marketing, selling, re-selling or distributing drugs, devices, services or therapies used to diagnose, treat, monitor, manage and alleviate health conditions.



Guide to Abstract Policies Governing ASCO Meetings

UPON SUBMISSION OF THE ABSTRACT TO ASCO

When submitting an abstract to any ASCO Meeting, the first author must agree to the following policies on behalf of all parties involved with the study. The first author is responsible for communicating the policies to all involved parties, including co-authors and sponsor, complying with these policies, and will be held accountable for any policy violations of the abstract.

- · Presentation Policy
- · Clinical Research Policy
- Copyright Transfer Policy

- Prior Presentation/Publication Policy
- · Confidentiality and Embargo Policy

Any violation of these policies may subject the abstract to rejection or removal from the ASCO Meeting.

Presentation Policy

The presenting author must agree to present the abstract in person if it is selected for presentation at the Meeting and adhere to the following guidelines for presenting authors.

At least one coauthor on the abstract must be eligible to present. If the first author is employed by a company or holds ownership with a private company (see company definition below), an alternate presenter who does not have a relevant employment or ownership relationship must be named from the list of coauthors. This applies to abstracts presented in an oral abstract session, rapid oral abstract session, or clinical science symposium. These presenters will also be subject to the same disclosure review and management strategies as faculty, per ASCO's Implementation Plan to Manage Relationships with Companies for CE Activities.

Company is defined as an entity whose business is developing, producing, marketing, selling, re-selling or distributing drugs, devices, services or therapies used to diagnose, treat, monitor, manage and alleviate health conditions.

Clinical Research Policy

Abstracts reporting on clinical research must verify that the work represented in the abstract was approved by an appropriate ethics committee or institutional review board and, if appropriate to this research, informed consent was obtained for all subjects.

Copyright Transfer Policy

The first author/presenting author must agree to:

- 1. Assign all copyrights of the abstract in all forms and media to ASCO effective if and when it is accepted for publication by ASCO.
- 2. Confirm that the abstract contains no material the publication of which violates any copyright or other personal or property right of any person or entity, and acknowledge that ASCO is relying on this representation in publishing this abstract.
- 3. Have obtained consent of all other authors to transfer copyright on their behalf, and indemnifies ASCO for any breach of this representation.
- 4. In the case of a "work made for hire" (a work prepared by an employee within the scope of his or her employment or commissioned as a work for hire under a written agreement), an authorized representative of the copyright owner must agree to the copyright transfer to ASCO. The submitting author must obtain consent of the copyright owner, and indemnifies ASCO for any breach of this representation.

--OR--

U.S. Federal Employment: If all named authors are officers or employees of the U.S. and the abstract was written as part of his or her official duties, it is not subject to U.S. copyright. Authors must also agree to items 2, 3, and 4 above. If such abstract is not published by ASCO, this form will not take effect.



Prior Presentation/Publication Policy

Meeting Presentations/Journal and Other Publications

Once submitted to ASCO and prior to the Meeting, the contents and conclusions of the abstract must not be presented at or published in conjunction with any scientific, medical, or educational meeting or published in any scientific, medical, or educational publication (in whole or in part). During the submission process, submitters are required to disclose if any part of the abstract has been previously presented or published and detail the updated data, results, and conclusions in their submission.

Exceptions

Studies previously accepted and presented at ASCO Meetings are eligible for resubmission and presentation at any other ASCO Meeting, including the ASCO Gastrointestinal Cancers Symposium, ASCO Genitourinary Cancers Symposium, ASCO

Quality Care Symposium, and ASCO Breakthrough. Authors are strongly encouraged to provide updated data in the abstract, as the novelty of the data will be considered during the abstract peer-review and selection process.

Previously presented or published Trials in Progress (TPS) abstracts are eligible for submission; however, if abstract copyright has already been assigned to another entity, the first author must obtain permission from that entity for ASCO to reprint the abstract. Permissions should be sent to abstracts@asco.org.

Preprints (non-peer-reviewed online comment drafts) are permitted until the abstract is submitted and may not be updated after submission.

FROM SUBMISSION OF ABSTRACT UNTIL ASCO PUBLISHES THE ABSTRACT

Confidentiality and Embargo Policy

Abstracts submitted to any ASCO Meeting are confidential and embargoed until ASCO publishes the abstract online. Latebreaking abstracts are confidential and embargoed once the study data is submitted to ASCO. For data previously presented or published, the Policy applies only to new or updated data in the abstract.

After the abstract is submitted to and prior to the abstract information being published online in conjunction with an ASCO Meeting, the author, coauthors, sponsor of the research, journalists, and others must not:

- make the information public or provide it to others who may make it public (such as news media), or
- · use the information for trading in the securities of any issuer or provide it to others who may use it for securities trading purposes.

Exceptions

Top-level study results may be presented and discussed at closed (nonpublic) meetings, such as investigator or cooperative group meetings, so long as meeting information and materials are not publicly discussed or disseminated, including on social media or online. Review the Frequently Asked Questions available on am.asco.org for more information.

ASCO recognizes that certain federal and international laws require disclosure of certain clinical trial results 1) through federal and international registries within a specific time period of trial completion, or 2) in relation to required disclosure by federal and international agencies for regulatory purposes related to drug safety and efficacy. Should disclosure of confidential information be required in either of these circumstances before ASCO publishes the abstract online, the required disclosure will not be viewed as a breach of ASCO's Embargo Policy.

Other than the purposes outlined above, exceptions to ASCO's Confidentiality and Embargo Policy require at least 48 hours advance communication with ASCO prior to any public release. Review ASCO's guidance and information for Reguesting an Exception listed further below on this page and direct all communications to CPexceptions@asco.org.

Even when an exception applies or is granted, ASCO retains the right, in its discretion, to accept or not accept any abstract for the ASCO Meeting on the basis of peer review, and once an abstract is accepted, to place the abstract or change its placement in the ASCO Meeting program depending on the extent of information released. If an exception applies or is granted, the study is unlikely to be included in the official press program for the ASCO Meeting.

Announcing Acceptance of the Abstract

Once an abstract has been officially accepted for presentation/ publication as part of the Meeting, authors and study sponsors are welcome to publicize the abstract's acceptance. However, the abstract title or session/presentation information should

not be shared until ASCO has publicly released this information. No study data or results can be included in this type of announcement (tweet, press release, etc.). Review the Frequently Asked Questions available on am.asco.org for more information.

FROM ABSTRACT ONLINE PUBLICATION UNTIL MEETING **PRESENTATION**

Once ASCO publishes the abstract online in conjunction with the ASCO Meeting, the general study findings may be discussed/ reported publicly. However, formal publication in a journal and release of full presentation slides or poster is prohibited until 7:00 AM (CT)/8:00 AM (ET) on the day of presentation. For publication-only abstracts, journal publication and release can occur at the time of abstract publication. Press releases issued after ASCO's online publication do not violate ASCO Policies.

View ASCO's abstract publication schedule and meeting program for abstract publication and presentation times.

For questions regarding resubmission of your ASCO abstract to a subsequent Meeting, consult the Policy for Resubmission of ASCO Abstracts to Other Meetings below and contact permissions@asco.org.

ADDITIONAL GUIDANCE TO REQUEST AN EXCEPTION TO ASCO'S CONFIDENTIALITY AND EMBARGO POLICY

Other than required disclosure for regulatory purposes as outlined above, exceptions to the Confidentiality and Embargo Policy may be granted by ASCO in extremely rare circumstances for public health reasons or to meet the requirements of state, national, or international government agencies. In these rare cases, requests should be directed to CPexceptions@asco.org with at least 48 hours notice for step-by-step guidance.

SEC Exceptions

A publicly traded company may determine that it is legally required to disclose certain data or other information from a confidential abstract in advance of the public release date to satisfy requirements of the U.S. Securities and Exchange Commission or a corresponding regulatory body in a country where the company's stock is traded (collectively, "SEC"). This need typically arises when there is a substantial likelihood that the information would be considered material by a reasonable investor, significantly altering the total mix of information available to the investor.

In general, an abstract in this situation is still eligible for inclusion in the ASCO Meeting provided that the company submits to ASCO, in advance of any public release, a letter signed by the company's legal counsel that contains the abstract title, indicates the format/nature of the public disclosure, and attests that (a) public disclosure of the information is necessary for the company to comply with applicable securities laws, and (b) the information

disclosed is the minimum necessary for such compliance. In addition, ASCO requires that the lead author be copied on the company's request for an exception. If the submission is in order, the SEC exception is self-executing and does not require preapproval from ASCO. If an SEC exception applies, the abstract is eligible to be peer reviewed and will not be rejected or removed from the meeting on the basis of a Confidentiality and Embargo Policy violation.

To the extent that the SEC exception applies, corporate partners of the company may, jointly or separately, issue a press release with the same information at that time. The abstract itself may not be released publicly by the company or lead author, as ASCO holds the copyright to the abstract.

Subject always to the company's regulatory obligations, ASCO would strongly prefer that the company's press release:

- · summarize study data cited in the abstract in a qualitative fashion rather than providing specific quantitative information;
- · avoid interpretations about the implications of the data for practice; and
- note that full data has been submitted to the ASCO Meeting.

By way of illustrating these preferences, a statement that a study "met its primary endpoint of increasing survival" is considered qualitative, while a statement that "survival was increased by 20% with the study drug" is considered quantitative. A quote such

as "We are encouraged by these promising results" would not be viewed as interpretive, while a quote such as "These findings support this drug as first line therapy in lung cancer" could be seen as an interpretation of the data. Information that is also appropriate for a press release includes that which is already publicly available.

For companies' convenience, a sample press release is available further illustrating these preferences.

If the press release or press coverage conveys significantly more information than ASCO's stated preferences and illustrated by the sample press release, the abstract may or may not be accepted into the ASCO Meeting on the basis of peer review. If the abstract has already been accepted when the press release is issued, the abstract's placement in the meeting program may be changed.

The exception will be publicly noted on ASCO's website once the abstract has been formally accepted to the Meeting.

RESUBMISSION OF ASCO ABSTRACTS TO OTHER MEETINGS

The procedure for submission of abstracts that have been accepted by an ASCO meeting to another meeting is governed primarily by the organizing body of that other meeting and secondarily is subject to ASCO's copyright in that abstract. If the organizing body does accept previously published/ presented abstracts, then ASCO will allow authors to submit their abstracts to such organizing body subject to compliance with the guidelines below and in accordance with ASCO's Confidentiality and Embargo Policy. The abstract may not be published until after ASCO publishes the abstract nor may the meeting presentation take place until after presentation at the ASCO Meeting.

On submission of the abstract, authors must notify the organizers of the other meeting that the abstract was presented at an ASCO Meeting, and that ASCO holds the copyright to the abstract.

A credit line must be published with the re-publication of the abstract as follows: "© [YEAR OF MEETING] American Society of Clinical Oncology, Inc. Reused with permission. This abstract was accepted and previously presented at the [YEAR AND NAME OF MEETING]. All rights reserved."

For additional information on ASCO copyright and approvals for reuse, please email our permissions department at permissions@asco.org.



am.asco.org/abstract-policies-embargoes-exceptions



Abstract Review and Selection Process

All abstracts are reviewed and discussed by the ASCO Scientific Program Committee by blinded review. Additionally, in advance of reviewing all submitted abstracts, all members of the committee receive training on mitigating biases when conducting scientific peer review.

Abstracts of superior quality will be selected for presentation at the Annual Meeting and for publication in the 2025 ASCO Annual Meeting Abstracts, an online supplement to the Journal of Clinical Oncology.

Regular Abstracts and Late-Breaking Abstracts

Abstract submissions are considered for all types of presentation, and as such authors are not permitted to state a preference for presentation type at the time of submission. Abstracts will be judged solely on the data submitted. Statements such as "further data will be presented" are not acceptable and will decrease the likelihood that the abstract will be selected for presentation at the Annual Meeting.

Trials in Progress Abstracts

Abstracts are considered for poster presentation and evaluated on the following criteria:

- · Strength of Science: Does the trial address an important and novel question?
- · Trial Design: Are the eligibility criteria, study endpoints, and planned analysis well defined in this abstract?
- Collaboration: Is there potential for investigator collaboration?
- Relevance: Will the results be relevant and of interest to the ASCO Annual Meeting community?
- · Requirements:
 - · Trial is registered, open, and enrolling patients.
 - Abstract does NOT contain preliminary data or results.

Abstract Notifications

Each first author will receive an email acknowledging receipt of the abstract after initiating a submission and after completing a submission. The first author will receive a letter of notification from the Scientific Program Committee regarding its decision by March 25, 2025.

Abstract Corrections and Withdrawals

The deadline to submit an abstract is January 28, 2025. All steps must be complete by the deadline, including adding all authors and their disclosure information. The designated submitter or the first author may view and modify their abstracts before the submission deadline. Your ASCO username and password are required to log in and edit the submission.

Corrections

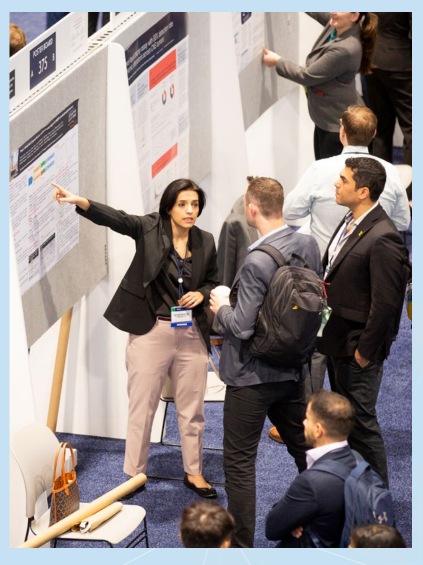
To protect the integrity of the review process, changes to abstract data or authors and disclosures will not be accepted after submission. Minor correction requests must be submitted by April 6, 2025, by the first author via email to abstracts@asco.org. These requests will be reviewed on a case-by-case basis.

After publication, corrections for major errors or omissions will be considered for six weeks after the meeting concludes. Approved changes will result in a formal erratum published in the Journal of Clinical Oncology.

All requests must be submitted by the first author. Requests from coauthors, pharmaceutical companies, or medical communication companies will not be accepted.

Withdrawals

Requests to withdraw an abstract from publication will be accepted through April 6. Requests received after this date will be reviewed on a case-by-case basis. All withdrawal requests must be made by the first author via email to abstracts@asco.org.





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- · Be the first author on the abstract submission and agree to present the abstract if selected for presentation at the Meeting.
- · Be enrolled in a fellowship/residency/advanced degree training program or in a medical or doctoral degree program at the time of abstract submission.
- · Provide a letter from their training program director or faculty advisor confirming the candidate is enrolled in the training program at the time of submission.
- · Provide a curriculum vitae.

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