SPECIAL REPORT

The International Collaboration for Cancer Classification and Research

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Abstract

Gaps in the translation of research findings to clinical management have been recognized for decades. They exist for the diagnosis as well as the management of cancer. The international standards for cancer diagnosis are contained within the World Health Organization (WHO) Classification of Tumours, published by the International Agency for Research on Cancer (IARC) and known worldwide as the WHO Blue Books. In addition to their relevance to individual patients, these volumes provide a valuable contribution to cancer research and surveillance, fulfilling an important role in scientific evidence synthesis and international standard setting. However, the multidimensional nature of cancer classification, the way in which the WHO Classification of Tumours is constructed, and the scientific information overload in the field pose important challenges for the translation of research findings to tumour classification and hence cancer diagnosis. To help address these challenges, we have established the International Collaboration for Cancer Classification and Research (IC^3^R) to provide a forum for the coordination of efforts in data comparability, standard setting and best practice recommendations in the field of tumour classification. The first IC^3^R meeting, held in Lyon, France, in February 2019, gathered representatives of major institutions involved in tumour classification and related fields to identify and discuss translational challenges in data comparability, standard setting,

List of Abbreviations: AI, artificial intelligence; AJCC, American Joint Committee on Cancer; ctDNA, circulating tumour DNA; ENCR, European Network of Cancer Registries; EQA, external quality assessment; EQUATOR, Enhancing the Quality and Transparency of Health Research; FIGO, International Federation of Gynaecology and Obstetrics; GA4GH, Global Alliance for Genomics and Health; GenQA, Genomics Quality Assessment; ClinGen, Clinical Genome Resource Somatic Working Group; GNU, GNU’s Not Unix; GPU, graphics processing unit; HGVS, Human Genome Variation Society; IARC, International Agency for Research on Cancer; ICCR, International Collaboration on Cancer Reporting; IC^3^R, The International Collaboration for Cancer Classification and Research; ICD-O, International Classification of Diseases for Oncology; ICD-10/11, International Classification of Diseases; IQN Path, Quality Network for Pathology; ISO, International Organization for Standardization; MVLD, minimum variant-level data; PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-Analyses; PT, proficiency testing; R&D, research and development; SEER, Surveillance, Epidemiology, and End Results; SNOMED CT, Systematized Nomenclature of Medicine Clinical Terms; STARD, Standards for Reporting Diagnostic Accuracy Studies; TNM, Tumour, node, metastasis; TRIPOD, Transparent Reporting of a Multivariable Prediction Model for Individual Prognosis or Diagnosis; UICC, Union for International Cancer Control; VCF, variant call format; WHO, World Health Organization.

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quality management, evidence evaluation and copyright, as well as to develop a collaborative plan for addressing these challenges.

KEYWORDS
cancer research, evidence-based pathology, International Agency for Research on Cancer (IARC), international standards, WHO Tumour Classification

1 | BACKGROUND

Cancer research aims to help people who are at risk of (or who already have) the various forms of cancer, in terms of risk assessment, prevention, early detection, diagnosis and treatment. Most countries have clear pathways for the translation of results of clinical trials of interventions into practice through drug approval and clinical guidelines, and also regulatory control over the use of medical devices. However, diagnostic interpretation, early detection procedures and screening methodologies are often not subjected to the same level of scrutiny.

The primary route for the incorporation of noncommercial knowledge into patient management is through the development of international standards and clinical guidance. The World Health Organization (WHO) Classification of Tumours fulfils an important role for cancer-related research and practice by providing an international consensus on classification and diagnostic criteria of tumours. The classification comprises a unique synthesis of scientific evidence (both written and illustrated) that underpins the diagnosis of individual tumour types. It is published as a series of books, known to pathologists as the WHO Blue Books, and is now also available as a website (see https://tumourclassification.iarc.who.int). The evidence base for each tumour type comprises peer-reviewed publications in multiple fields, and although the primary diagnostic method is still histopathology, other disciplines have an increasingly large role to play. The vast number of publications that inform the classification now present a significant problem of information overload, but also an enormous opportunity, as our understanding of cancer continues to progress. It is also true that not all of the information published warrants inclusion in the WHO classification or in clinical guidance—it is of variable quality and relevance, and sifting through this information is now a major task for the biomedical community. Systematic reviews, which have been the bedrock of evidence-based medicine during the last decades, have proven to provide reliable evidence synthesis and to allow the evaluation of available information (eg, the application of such methods, in addition to the attachment to existing guidelines as the EQUATOR (Enhancing the Quality and Transparency of Health Research) Network guidance (see https://www.equator-network.org/) would certainly help to get results into practice in the field of cancer research and tumour classification. But focus on translational research is also needed to bridge the gap between basic and clinical investigators, and facilitate the application of research results to practice. The task is still enormous and needs individuals, organizations and networks committed to bridge the gap.

What’s new?
The World Health Organization Classification of Tumours has been informing cancer research and clinical practice by providing an international consensus on the tumour criteria for cancer diagnosis. In a new initiative coordinated by the International Agency for Research on Cancer, major institutions are now joining forces to address the remaining challenges in translational research and facilitate the application of research results to clinical practice. The newly-established International Collaboration for Cancer Classification and Research (IC²R) aims to provide a forum for the coordination of efforts in generating evidence, setting standards, and providing best practice recommendations for tumor classification and cancer research.

1.1 | Vision and aims

IC²R will promote collaboration between organizations to ensure and improve the provision of high-quality evidence and evidence
synthesis, permitting rapid translation of cancer research into clinical diagnostic practice for patients.

IC3R will be tasked with the following:

- Harmonizing cancer-related data
- Setting standards for analytical procedures
- Identifying and informing the community of critical gaps (e.g., non-uniform annotations and classifications and gaps in bioinformatics, computational pathology, clinical chemistry and other areas of research)
- Managing quality and establishing best practice
- Coordination of developments in artificial intelligence (AI)
- Updating the evidence and generating evidence synthesis
- Addressing copyright considerations

IC3R will act in close collaboration with other stakeholders in the field. The International Agency for Research on Cancer (IARC) will provide the secretariat and may assist the coordination of international datasets as required, as well as encouraging the generation of standards and procedures for data management that will benefit all parties involved.

The activities of IC3R currently fall into six main categories (although other activities could be added if members identify the need): harmonizing data, identifying gaps, setting standards, improving quality and establishing best practices, updating the evidence, and addressing copyright considerations. In our study, we provide a short description of these categories, the challenges and potential solutions IC3R will focus on.

2 | HARMONIZING CANCER-RELATED DATA

Tumour classification relies on various types of data related to the patient, clinical context, biological specimens and analytical measurements. Harmonizing the structure, collection and registration of data across cases within an institution is essential to ensure comparability and reproducibility, but standardization and harmonization across institutions is even more crucial, to improve the basis of tumour classification and inform cancer care worldwide.

The need for data standardization and harmonization is present at all stages of the tumour classification process. For example, cancer registration standards for data collection are needed to provide alignment for the determination of the date of diagnosis between groups such as the Surveillance, Epidemiology and End Results (SEER) program, the International Association of Cancer Registries (IACR) and the European Network of Cancer Registries (ENCN). Consent on criteria for the definition of case entities is important, as for example in patients with multiple primary tumours. Differences between the TNM (tumour, node, metastasis) staging systems of the American Joint Committee on Cancer (AJCC) and the Union for International Cancer Control (UICC), and between TNM and International Federation of Gynaecology and Obstetrics (FIGO) staging are increasingly addressed collectively by these organizations and benefit both patients and research. Standardization is also needed in the pipelines used for the identification of relevant genomic characteristics and the reporting of molecular biomarker assays, where insufficient laboratory certification, absence of standards and lack of confirmatory large-scale studies for the validation of biomarkers often limits the scope of the results. Some current initiatives aim to address these issues, but do not cover all eventualities, for instance, the International Collaboration on Cancer Reporting (ICCR), a collaborative initiative that involves key international cancer- and pathology-related organizations. ICCR produces internationally standardized and evidence-based datasets for the pathology reporting of cancer using defined terminology and normalized pathology information on tumour classification and staging, as well as on prognostic and predictive factors (see http://www.iccr-cancer.org/). These datasets are produced in a variety of formats and languages to maximize their worldwide usage.
TABLE 1  Online databases for molecular profiling of tumour DNA, RNA and epigenetic features

| Name                                                                 | Available at                                                                 |
|----------------------------------------------------------------------|-------------------------------------------------------------------------------|
| The Catalogue of Somatic Mutations in Cancer (COSMIC)                | https://cancer.sanger.ac.uk/cosmic                                            |
| The National Cancer Institute's Genomic Data Commons (NCI GDC)       | https://gdc.cancer.gov/                                                        |
| The International Cancer Genome Consortium (ICGC)                    | https://icgc.org/                                                             |
| The cBioPortal for Cancer Genomics                                   | https://www.cbioportal.org/                                                   |
| The National Center for Biotechnology Information's Gene Expression Omnibus (NCBI GEO) | https://www.ncbi.nlm.nih.gov/geo/                                                |
| The European Bioinformatics Institute's ArrayExpress Archive of Functional Genomics Data (EBI ArrayExpress) | https://www.ebi.ac.uk/arrayexpress/                                           |

Large-scale molecular profiling of tumour DNA, RNA, and epigenetic features has generated informative data for tumour classification. These data are available through various online databases (see Table 1), which have each implemented their own data harmonization procedures.\textsuperscript{4-6} Also, for the reporting of genetic variations well-accepted standards exist, the Variant Call Format (VCF) and the Human Genome Variation Society (HGVS) nomenclature, which have been adopted in a growing number of resources. However, the use of variations of these formats impedes the achievement of clear equivalence between data sources and engenders difficulty in the comparability of research results.

Many current genomic resources suffer from a lack of harmonization of molecular data, clinical information and biological specimen features. Few of them use recognized international classifications such as the International Classification of Diseases for Oncology (ICD-O), International Classification of Diseases (ICD-10/11) or Systematized Nomenclature of Medicine Clinical Terms (SNOMED CT). Many have difficulties due to the diversity of bioinformatics pipelines used for the processing of genomic data, which may create batch effects when aggregating data from different databases to conduct large-scale studies. Here too, some initiatives to address these issues are ongoing; an example is the proposed guidelines for the harmonization of variant calling across human genetics projects.\textsuperscript{7,8} The Global Alliance for Genomics and Health (GA4GH) was set up to create frameworks and standards to enable the responsible, voluntary and secure sharing of genomic and health-related data, and the Clinical Genome Resource (ClinGen) Somatic Working Group has developed a minimum variant-level data (MVLD) methodology for promoting the harmonization of cancer variant representation and efficient distribution of this information through ClinVar.\textsuperscript{9} Beyond standards of nomenclature, the interpretation of the significance of the specific variants in terms of their clinical impact is evolving and improving, but many variants still fall into the challenging category of “variants of unknown significance”. The development of a standardized terminology and dataset for interoperability would accelerate the challenging process of determining the clinical significance of specific gene variants.

These initiatives pave the way for improving database interoperability and facilitating the aggregation of datasets, but important challenges remain in the harmonization of individual data and production of datasets usable for evidence-based medicine. Data quality metrics and assessments should also be standardized, and internationally approved guidelines should be developed with recommendations for minimum data sets, content and validation, as well as reporting recommendations defining the minimum annotations needed, and quality standards for cancer research databases and tumour classification.

3  IDENTIFYING GAPS

The WHO Classification of Tumours is a taxonomy. It groups tumours into diagnostic entities (tumour types) based on their shared characteristics, just as Linnaeus (1758) identified species based on shared characteristics. There is still a reliance on histology and increasingly on molecular pathology, but radiology, epidemiology and other disciplines contribute to the classification of many tumour types. The classification defines a tumour type as an entity in which multiple parameters (eg, radiological appearance, localization, histopathology, and molecular profile) differ from those of other types, whereas a subtype may not vary by more than one or two parameters, and usually without sufficient evidence that these differences would currently affect the clinical management of the patient.

The WHO Classification of Tumours also identifies tumour types or subtypes where there is currently insufficient information about some characteristics (parameters), but although this lack of information may be indicated in the WHO Blue Books, there is no organized database that systematizes the identified research needs in a way that could drive research to bridge these gaps. As part of IC\textsuperscript{3}R, IARC proposes to maintain an organized record of the current research needs, with contributions from other organizations.

As an example of an as-yet unmet research need, one of the challenges of personalized medicine is to identify the specific molecular anomalies that define tumour types and subtypes. The WHO Blue Books focus on incorporating the most frequent anomalies, but these studies also provide a steady flow of newly described rarer molecular features that identify tumour types that are not necessarily published, due to insufficient information and validation of these characteristics and their relevance. Similarly, as technologies evolve and the research community applies them in innovative ways, the information about the molecular features used in tumour classification becomes increasingly diverse and complex. For these reasons, evaluations of the existing evidence for mutations and their relationships to clinical features such as diagnosis and prognosis are required for each tumour type and recommendations to guide future research in the production of missing
evidence need to be formulated. This information must be reliable and reproducible to permit informed, evidence-based decisions.

The effort to identify relevant anomalies cannot be fuelled solely by case reports since they have a high risk of bias in their observations due to limitations of the study design. Publication policies often lead editors to accept the first report of a new anomaly, but further identical reports are viewed as less interesting in terms of novelty and the submitted paper can be rejected (an example of publication bias). The establishment of open and interactive database(s) that describe all relevant features and their clinical or pathological relevance is to be encouraged, though a multiplicity of such databases can result in a further need for harmonization (eg, Decipher; https://decipher.sanger.ac.uk). IC3R community could stimulate the development of a forum through which additional knowledge gaps could also be communicated to these databases, which would facilitate the evaluation of the available evidence for the associations of these features with relevant clinical outcomes. The resulting technical developments and research could subsequently be used to inform the WHO Classification of Tumours. Similarly, this community may also provide the potential to build further consortia to provide the research community with the biological and technical infrastructure necessary to validate and replicate key insights, as well as to communicate cutting-edge developments (eg, future AI-based analysis or other integrated analysis approaches). Such a database, and the related community, would be a valuable addition to facilitating an appropriate translation of robust findings from the research community into practice.

4 | SETTING STANDARDS

The rapid, global development of the healthcare industry and related research has meant that increasingly greater importance is being placed on standardization activities that facilitate compatibility. Standards for technology ensure that any expected component or process will operate safely and predictably, independently of the manufacturer and/or operator. The same considerations apply to the standardization of processes and services. Service standardization in health care is an important adjunct to promoting the provision of high-quality, specialized expertise and services globally. Systems in several countries provide standards accreditation through national structures and maintain online platforms that monitor quality indicators for nationwide analysis.

In terms of current research in cancer, conclusions related to tumour aetiology often follow the law of large numbers, where increasingly larger sample sizes are used for analysis. Standards are essential to support such scalability in medical research. However, the level of standards implementation is not equally supported or reported across different geographical regions; as a result, the promotion, endorsement and development of such standards are often delegated to national and/or international professional associations that face limitations implementing and monitoring these standards. Stronger international commitment is required to promote the development and subsequent uniform implementation of standards in cancer research.

The Standards for Reporting Diagnostic Accuracy Studies (STARD) constitute a good example of such an initiative, which was led by the professional community, and is now part of the EQA

5 | MANAGING QUALITY AND ESTABLISHING BEST PRACTICE

5.1 | The need for quality improvement and management

Quality management systems are designed to plan, control and improve the elements of a process that impact on the achievement of the desired results, which includes the safety, effectiveness and accuracy of testing and has a fundamental role in diagnostic science. These systems permit cross comparison between diagnostic services. All cancer research laboratories, including pathology, genetics and other diagnostic services should implement minimum standards and follow the continuous model required by these systems. They also allow the continuous monitoring of performance indicators and the reporting of poor performance in cases where the required standards have not been reached. This gives laboratories the opportunity to investigate the causes and determinants, for instance performing a root cause analysis that would permit the planning of specific actions to rectify the problem and promote improvements. This type of process management ensures reliable test results and that accurate interpretations are provided to patients via the referring clinician. Many processes are performed in diagnostic services to guarantee the accuracy, reliability and traceability of results (eg, it is expected that the correct international nomenclature is used for an external quality assessment (EQA) in a genetics/molecular laboratory, and they often apply programs as EQA or proficiency testing (PT). It is also important to note that the application of an ISO-accredited EQA provides evidence of the standard of testing and reporting (including
accuracy, reliability and appropriateness) for diagnostic services, as well as benchmarks, and should be undertaken at least annually for every aspect of the service.

5.2 Methods of quality improvement

Quality management systems provide methods for centres to assess their clinical and analytical performance, as well as their interpretation of molecular pathology/genetic testing based on international standards and compared to that of other laboratories. Additionally, EQA programmes offered by external providers facilitate method validation, comparing of results among laboratories, problem identification and compliance with standards and requirements, as well as increases credibility.

The external audit of the diagnostic services is an essential part of any type of quality management programme and is usually undertaken by an EQA or PT provider and leads to the certification of procedures or accreditation of the services based on international ISO standards. These programs are excellent tools to keep procedures and all related variables (staff, equipment and method) well controlled, ensuring that the quality of the service meets minimum professional standards and continuous improvement principles are followed. EQA providers need to be autonomous from professional and national bodies to provide independent external verification of service quality, in addition to a performance guarantee to the laboratory, the host institution and service users. EQA providers have the ability to compare participating laboratories, thus benchmarking the best practices for services. These comparisons between laboratories are incredible helpful in refining laboratory standards and essential to establish best practice guidelines. This is feasible for research laboratories, and some already participate in clinical EQA schemes. To promote this practice at worldwide level, the cancer research and tumour classification community will enable to enhance collaborative projects and improve comparability of results in the field. IC3R constitutes an opportunity to appraise EQA programs in cancer research and promote international comparable evaluations.

5.3 Benefits of external quality assurance schemes

Among their many benefits (see Table 2), EQA programmes improve and validate the overall quality of a service. In the case of a diagnostic, genomic or genetic service, this includes all procedures from sample...
TABLE 2  Benefits of applying an external quality assessment

- Makes possible consistent quality assessment of diagnostic and/or clinical services.
- Permits the establishment of a continuous quality improvement plan.
- Allows identifying imperfect practices and errors, thus improving quality and ensuring patient safety.
- Identifies problems/flaws with diagnostic procedures and kits.
- Recognizes inefficiencies, improving the diagnostic service of laboratories.
- Assesses the laboratory’s internal quality control, equipment, training, and so on, that is, in the quality management program.
- Serves as an aid for centres to stay up to date.
- Ensures uniformity in practices and aids in the development of new ones.
- Permits to establish quality standards at national level and across borders.
- Possibilities for patient involvement with the aim of improving the overall service to the user.
- Provides external verification/validation of service quality and building public confidence.
- Playing a surveillance role: ensuring laboratories’ adherence to professional guidelines and international standards.
- Ensuring that the provision of genetic counselling is the same for all family members.

receipt to genetic referral, including necessary education and training. EQA allows the identification of variations in results and reporting. It encourages a closer interaction between laboratories and clinicians, and often underpins the development of guidelines to promote consistency in the procedures and ensure high-quality testing for the benefit of patients.

Several international entities offer EQA programs in the field of molecular pathology, and most are ISO 17043 accredited. Laboratories clearly benefit from selecting an ISO-accredited EQA provider that covers the procedures of their diagnostic service, but there is not always one available. Pilot EQAs are not eligible for accreditation since they have no performance monitoring, but after the first year of participation, it should be mandatory to include the EQAs in the Quality Management System for both research facilities and diagnostic services in cancer. Genomics Quality Assessment (GenQA) and other EQA providers have proven that such annual participation improves the quality of the diagnostic service.20 For instance, the recent EQA assessment of DNA extraction from formalin-fixed and paraffin-embedded samples by Professor Z. Deans from GenQA showed an improvement in DNA quality and quantity over time.24

A continuous effort is made by various EQA providers to standardize the procedures and the scoring system of the EQA schemes in molecular pathology21 so that a large number of diagnostic services can participate. To assist diagnostic and research centres, GenQA provides teaching and education through an online training and competency tool, G-TACT, for various aspects of a cancer genomics service.25 To ensure consistency among EQA providers and to develop new operating models, the most relevant EQA providers recently founded the International Quality Network for Pathology (IQN Path). IQNPath has the main purpose of improving the quality of biomarker tests, in particular with regard to molecular pathology.

Although current EQA schemes clearly demonstrate the challenges faced in cancer diagnostic testing and interpretation, as well as the need for better training, competency assessment and standardization worldwide, the results from assessments demonstrate the value of consistent data, sharing of results and collaboration in improving diagnostic practice.

5.4  |  Quality improvement and research

Improving quality in cancer-related research is of relevance for tumour classification and diagnostic practice in general.26 Applying lessons learned from quality management in laboratory medicine could drive improvement in the quality of cancer research and stimulate better translation of research findings into clinical practice. However, this is not always easy. A recent systematic review of the use of circulating tumour DNA (ctDNA) for early cancer detection reached the conclusion that several preanalytical, analytical and postanalytical considerations need to be addressed before biomarkers enter clinical practice.27 The same issue is shown by data from EQAs, suggesting that the quality of ctDNA testing needs to be significantly improved.28

The practice of quality management in cancer research laboratories could help to solve such problems. The adoption of ISO standards for cancer research laboratories, perhaps adapted from ISO 15189 for clinical laboratories, could provide a quality management system; training; an environmental policy; guidance for equipment, information systems and materials, as well as defined preanalytical, analytical and postanalytical processes (standard operating procedures); evaluation methods (including publication); and quality assurance. Good scientific practice already mandates much of this and many important research institutions have defined guidelines for effective validation of some procedures as next-generation sequencing methods and monitoring of analytical procedures.29-31 Additionally, the European Molecular Genetics Quality Network and GenQA have launched a pilot EQA methodology for next-generation sequencing in Europe.26,29,32-34 Thus, some research laboratories already participate in EQA schemes driven by the above initiatives, but consolidation of this practice is necessary to promote wider, international standardization.

In addition, a more rigorous attitude in reporting of results and scientific publication is desirable. For example, differences in sample collection, extraction, storage and processing (physical and computational) can have a strong effect on downstream analyses and decisions. Reporting of raw data in open-source databases is a desirable standard that facilitates pooled analyses and accelerates knowledge generation. Many scientific journals have already adopted the recommendations of the EQUATOR network for scientific publications,35 and ICRs will endorse these journals while at the time as it promotes the application of standards in the cancer research community.
6 | KEEPING ABREAST OF DEVELOPMENTS IN AI

In recent years, there has been a surge of interest and some significant developments in the emerging area of computational pathology, a rapidly developing discipline in its own right, concerned mainly with the computerized analysis of digitized images of histology slides. This surge of interest has been spurred by the increased use of digital slide scanners in diagnosis and significant advances in AI algorithms, in particular deep learning algorithms, matched by the availability of powerful graphics processing units (GPUs; originally developed for the gaming industry) and storage hardware at a fraction of the cost compared with that only a few years ago. In combination, computational pathology has been dubbed “the third revolution in pathology”, after immunohistochemistry and next-generation sequencing and has to be considered in IC²R's strategy.

6.1 | The rise of computational pathology

Synergistic technological developments in the areas of slide scanning, digital storage and high-performance computing have enabled the uptake of digital pathology for routine diagnostics, resulting in the generation of several terabytes of information-rich, high-resolution whole slide images from individual clinical histopathology departments (the so-called “big image data”) on a daily basis. The raw pixel information in whole slide images is ripe for AI-based automated or computer-assisted diagnosis and subtyping of cancer in a fast, accurate, objective and reproducible manner. Whole slide image pixel data with linked clinicopathological data are also an invaluable resource for mining and discovery of deep spatial patterns and novel image-based markers of direct benefit to patients in terms of prediction of recurrence, progression and response to therapy, paving the way for personalized treatment. Some of the notable developments in the area of computational pathology include the discovery of stromal nuclear features shown to be more strongly associated with survival of breast cancer than tumour cell features, the Immunoscore method for predicting outcome in Stage I colon cancer, superhuman performance of AI algorithms in the detection of lymph node metastasis in breast cancer, nuclear shape and orientation features for predicting oncotype DX-risk categories, and digital scoring of the abundance of tumour-infiltrating lymphocytes in oral cancers.

6.2 | Benchmark datasets for AI R&D and validation

Modern deep learning algorithms are known to be generally “data hungry”. These algorithms are also quite vulnerable to a data bias problem where assurance is needed that the dataset used for prediction is as fully representative as possible of the phenomenon to be predicted. In future, the construction of large multicentric benchmark datasets will be key to the research and development (R&D) of novel AI algorithms that are robust and reliable in addition to international standards. Such benchmark datasets can also serve as a vehicle for the rapid evaluation and validation of AI algorithms for regulatory approvals. With these objectives in mind and to achieve higher reproducibility in computational pathology, the PathLAKE (Pathology Image Data Lake for Analytics Knowledge & Education) Centre of Excellence in the United Kingdom is putting together a massive repository of multicentric pathology images linked with diagnostics and clinical outcome data.

6.3 | AI-based evidence synthesis

In recent years, AI algorithms have demonstrated significant potential for synthesis of evidence—one of the key objectives of IC²R—in particular, tools for identifying randomized controlled trials and assessment of bias in their reporting. These tools are still in the early stages of development and will need to be rigorously validated. Nevertheless, they can be used to assist with systematic reviews for evidence synthesis, saving time and costs in the synthesis. It will also be of paramount importance to ensure that key principles of evidence synthesis (eg, inclusivity, transparency, rigor and accessibility, as recommended by the UK Royal Society and Academy of Medical Sciences), are upheld when AI algorithms are deployed for the purposes of evidence synthesis.

6.4 | Deep and integrative mining for better classification

AI-based deep and integrative mining of vast amounts of raw pixel information, together with linked clinical outcomes and other types of linked data (eg, genomic and physiological data), can also be used to revisit the existing taxonomies. In the genomic space, AI algorithms can be deployed for the purposes of evidence synthesis. As a specific example, there are >30 different subtypes of salivary gland tumours in the current WHO classification. However, the classification may not be of enough prognostic relevance or sufficiently informative on its own to drive patient management. AI-based integrated analysis has the potential to drill down into the data, pick out deep subvisual and integrative patterns, and form disease subgroups directly related to outcome. Although there are clear benefits to the detailed subtyping of tumours (such as avoiding suboptimal treatment of patients due to excessively broad classifications), it would perhaps be better to have fewer subtypes that are more useful in determining the best course of action for a given patient than an overly detailed subtyping that is not as informative for patient management.

7 | UPDATING THE EVIDENCE

The available evidence must be reviewed periodically in order to keep tumour classifications up to date, but it is not always clear how or
when such reviews should be performed. Literature reviews produce variable results depending on the expertise and perspective of the reviewer and the methodology used, with the potential for relevant studies to be missed and interpretations of results to be misleading. In addition, not every statement needs to be informed by complex systematic reviews. Often, traditional background or scoping reviews provide sufficient information to evaluate a subject, and available resources can be saved to be used to inform pressing issues of tumour classification—these topics need to be assessed by systematic reviews due to their potential controversy. Evidence-based medicine has been providing recommendations and promoting the performance of systematic reviews as the cornerstone of its methods, encouraging comprehensive literature searching, transparency in methods and rigorous study appraisal. However, best practice guidelines, such as the Cochrane Handbook, and related guidelines, such as PRISMA, are closely aligned with meta-analytic reviews for medical interventions evaluated by randomized controlled trials—methods that are not necessarily appropriate for pathology. Evidence-based medicine principles and practices need to be adapted to be useful in tumour classification, and an evidence-based pathology approach that adjusts methods, provides recommendations and offers training is greatly needed.

There are additional challenges for an evidence-based pathology approach to the classification of tumours. Evidence levels and study design considerations are applied with great variations within the field, and if we consider the known evidence hierarchy, most pathology studies examine associations with the lowest levels of evidence. These studies usually do not consider the limitations of their study design and the consequent risk of bias needing to be considered in drawing conclusions and planning research. Pathology as a specialty would benefit from adding research methodology to the curriculum and continuous professional development for its specialists. Methodological discussions are also needed, as well as an initiative for expert consensus on principles and best practices in the field that would allow more consistent interpretation and comparison of studies in pathology. This would also identify areas where new solutions and underpinning methodological research are required.

Thus, evidence-based pathology will have to deal with the need for timely assessments of controversial decisions, the exponential rise in the number of scientific publications and the management of new types of information such as evidence from genetics or big data sources in bioinformatics. How much subgrouping of uncommon tumours can be sustained before the size of the groups worldwide becomes too small to prove that differences in behaviour or outcome could have occurred by chance alone? To address these challenges, new methodologies will have to be developed, a collective of skilled experts trained and an extended network of evidence-based pathology hubs created. Such an evidence-based pathology movement, with sufficient resources and adequate coordination, could respond with evidence-based assessments to the predicted large number of decisions that will have to be addressed in the near future for tumour classification as the number of potential classifiers continues to increase rapidly. IC^3R is ideally positioned to efficiently promote and coordinate such an endeavour, working with other partners in pathology and drawing on the expertise of the more widespread evidence-based movements.

8 | COPYRIGHT AND OPEN SCIENCE

Copyright law is somewhat at variance with the principles of open science. The restrictions that certain copyright transfer provisions impose on scientists remain a hindrance to information dissemination. However, this amounts to securing profit from work financed by others—often by public funds—and it delays the potential impact of translating discoveries into patient care. Open science is a movement that aims to make scientific research and data available to all, from professionals to citizens. Open science advocates for practices such as publishing open research and open access to publications in order to facilitate scientific growth, with the ultimate aim of making it easier to publish and communicate scientific knowledge. However, the transition towards effective open science is slow and challenging, and although journals do offer authors the option of submitting articles as open access, this is often accompanied by increased financial burden on the authors.

Open access is not an infringement on copyright, because authors or their institutions still own the original copyright for their publications. The problem is that they are often asked to transfer these rights to the publisher, allowing publishers to set the terms for providing open access. Ideally, an open license would be used to publish, with clearly defined access and reuse rights, and there are certain journals that do offer the option of publishing under a Creative Commons license (eg, CC BY 4.0 for publications and CCO for data) or GNU (for software and code). In addition, some large institutions, such as the National Institutes of Health (NIH), impose open access requirements for articles arising from their funding, requiring that the articles be made publicly available on PubMed Central. Moreover, movements such as OpenAIRE encourage researchers to choose publishers who allow them to retain their authors’ rights, attempting to change practices that limit knowledge exchange. Working to improve access to information, through various means, is something that IC^3R is committed to examining.

9 | CONCLUSIONS

The IC^3R has been founded to address the need of coordination and collaboration in the multidimensional environment of cancer research. The collaboration will provide essential guidance for involved parties, set standards in the field and provide evidence and evidence synthesis considering priorities in tumour classification.

The IC^3R work plan includes five main actions that will ensure the future of the consortium and the implementation of appropriate measures to meet the objectives (see supplementary material):
1. The establishment of a stable managerial structure to facilitate the planning and production of standards well into the future, as well as to design, coordinate and fund projects that address the identified challenges (harmonization of standards, information system development, evidence-based data, etc.) at the initiative of a steering group.

**Action:** A consortium has been founded to provide the managerial structure needed to facilitate the design and implementation of appropriate measures to meet the objectives detailed in our study. In the initial year, the structure and working organigram will be set up, including the coordination team (composed by a project manager and assistant part of the secretariat) and a communications strategy. In parallel, workgroups will be created to address the identified challenges (harmonization of standards, information system development, evidence-based data, etc.) at the initiative of the Steering Group.

2. Drafting of recommendations for quality management in research laboratories are one of the most pressing needs. Options for formal accreditations and best practices need to be provided.

**Action:** The adoption of an ISO standard for cancer research laboratories is in preparation.

3. Promotion and development of evidence-based pathology to provide the methodological and technical support needed in the field is also relevant and should be initiated as soon as possible.

**Action:** A project for the promotion and development of evidence-based pathology will be launched that will include deliverables such as evidence levels adapted for the field of Pathology, new systematic review methods, training in evidence-based pathology, adapted systematic review tools and a network of systematic reviewers (Figure 3).

4. Identification of tumour types for which there is insufficient information about their characteristics are clearly stated within the WHO Blue Books, but there is no organized database of the research needs identified or coordinated response to these needs. As part of IC3R, IARC should maintain an organized record of current research needs and allow other organizations to add to it.

**Action:** A database of current research needs will be established based on IARC publications and others, including a proposal for the prioritization of issues.

One of the key advancements provided by such a consortium and its activities would be the establishment of an open and interactive network of experts that describe for tumour classification relevant features, participate in the identification of needs and assess trends. Importantly, one might envisage the development of a community around this network that participates actively in the continuous improvement of the initiative.

**ACKNOWLEDGEMENTS**

We are grateful to Dr Louise Jones and Professor Mark Caulfield for their participation in the preliminary draft of the manuscript, to Asiedua Asante for her assessment and to Jessica Cox and Catarina Marques for language and image editing.
CONFLICT OF INTEREST
Dr Nicola Normanno reports grants and/or personal fees from Amgen, MSD, Qiagen, Roche, BMS, Merck, Thermofisher, Astrazeneca, Zanofi, ELI Lilly, Archer Dx and Illumina outside the submitted work and being President of the International Quality Network for Pathology (IQN Path) and the Italian Cancer Society (SIC). Dr Richard L. Schilsky reports his institution (ASCO) receives funding on his behalf from Astra-Zeneca, Bayer, Boehringer-Ingelheim, Bristol Myers Squibb, Eli Lilly & Co., Genentech, Merck, Pfizer. Dr Stefan Hold-enrieder reports grants from Roche Diagnostics and Volition RX, as well as being a Scientific Advisory Board Member and consultant of Volition RX. The other authors declare that they have no conflict of interest. The other authors indicated no financial relationships.

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The content of this article represents the personal views of the authors and does not necessarily represent the views of the authors’ employers or associated institutions. Where authors are identified as personnel of the International Agency for Research on Cancer/World Health Organization, the authors alone are responsible for the views expressed in this article, and these views do not necessarily represent the decisions, policy or views of the International Agency for Research on Cancer/World Health Organization.

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