Computational pathology in cancer diagnosis, prognosis, and prediction – present day and prospects

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Abstract

Computational pathology refers to applying deep learning techniques and algorithms to analyse and interpret histopathology images. Advances in artificial intelligence (AI) have led to an explosion in innovation in computational pathology, ranging from the prospect of automation of routine diagnostic tasks to the discovery of new prognostic and predictive biomarkers from tissue morphology. Despite the promising potential of computational pathology, its integration in clinical settings has been limited by a range of obstacles including operational, technical, regulatory, ethical, financial, and cultural challenges. Here, we focus on the pathologists' perspective of computational pathology: we map its current translational research landscape, evaluate its clinical utility, and address the more common challenges slowing clinical adoption and implementation. We conclude by describing contemporary approaches to drive forward these techniques.

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Introduction

Rapid advances in molecular diagnostics, bioinformatics, and computer hardware have facilitated innovation in healthcare technology, including improved diagnostic, predictive, and prognostic pathways and treatments for cancer patients [1]. With a globally ageing population and increased early detection, and thus increased incidence, of disease [2–4], the necessity for more costeffective, agile, efficient, and equitable healthcare systems is driving public and private funding streams in translational and commercial research in this space [5]. Diagnostic pathology, the gold standard of cancer diagnostics, is transitioning to a digital workflow (digital pathology) and will switch from bright-field microscopy to digital image assessment (Figure 1). This progression is spurring interest in computational pathology [6,7].

Over the last decade, progress in radiomics and multiomics technology has generated a comprehensive map of the molecular dynamics that govern cancer, aiding in the discovery of new prognostic biomarkers and treatment targets. Some technologies have successfully been

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Figure 1. Schematic diagram of the topics discussed in the review. In the conventional histopathology workflow, slides are assessed by a pathologist under a bright-field microscope. In the digital pathology workflow, slides are scanned with a whole slide image (WSI) scanner and pathologists assess WSIs digitally on a computer. Challenges to transit from a conventional to a digital pathology workflow are listed. Opportunities in the translational landscape and key topics for the adoption of computational pathology are indicated. This figure was created with the aid of BioRender.com.

translated to provide clinically usable insights and guide the care of cancer patients [8]. In contrast, conventional histopathology, at the core of cancer diagnostics, has yet to experience such a technological evolution [1,8]. Histopathologists not only diagnose cancer based on an assessment of tissue sections but also provide prognostic information, for example by subtyping and grading the disease, and patient treatment is largely decided on such characteristics. Decisions are based on complex visual features and require significant skill and training [9]. However, a shortage of new entrants in histopathology, along with an increasing workload, is of concern [10–12]. To tackle these challenges, advances must be targeted towards developing computational tools that can aid pathologists in performing their routine diagnostic work and alleviate their heavy workload, as well as potentially unlocking new insights to further precision medicine. Investments are needed in the following four critical computational pathology domains: the development of AI algorithms, the infrastructure and data management, the integration of machine learning and analytics into the diagnostic workflows, and collaborative research initiatives.

Computational pathology utilises deep learning methods to examine whole slide image (WSI) tissue samples. It aims to create algorithms to execute standard diagnostic procedures, to establish models that scrutinise tissue morphology to predict diagnostic and molecular changes or discover novel biomarkers [1,7,8,13,14]. However, there has been a slow uptake in the diagnostic setting due to several operational, technical, regulatory, ethical, financial, and cultural challenges. Centred on the pathologists' view, this review describes in detail the current computational pathology landscape, outlines key challenges hindering its progress, illustrates the commercial environment, and explores potential avenues towards adopting these technologies by healthcare systems (Figure 1).

The translational landscape

Routine histopathology diagnostics: cancer diagnosis, subtyping, and grading

The 2016 CAMELYON challenge was a launch pad for computational pathology research that stimulated collaboration between oncology, pathology, and computer science communities [15]. Both CAMELYON16 and CAMELYON17 challenges demonstrated AI's potential to automate the time-consuming diagnostic histopathological task of lymph node assessment for cancer staging purposes; convolutional neural network (CNN) models achieved comparable accuracies to pathologists in detecting metastasis [15–17]. Computational pathology algorithms based on CNNs have since demonstrated expert-level performance across a range of pathology tasks prone to inter-observer variability, including, but not limited to, diagnosing and grading prostate cancer [18–20], counting mitoses in breast cancer [21], grading tumour budding in colorectal cancer [22], or diagnosing, subtyping, and detecting associated gene mutations in lung cancer [23-27].

Detecting known biomarkers with computational pathology

pathology Computational methods have been implemented to capture known biomarkers directly from WSIs, such as the expression of oestrogen receptor (ER), human epidermal growth factor receptor 2 (HER2), and programmed cell death protein 1 (PD-1) and its ligand PD-L1 [27–29]. However, immunohistochemical (IHC) assays notoriously suffer from a lack of standardisation in both pre-analytical and scoring methodologies. Genetic alterations, on the other hand, may cause a diverse spectrum of morphological features, as is the case in salivary gland tumours where single gene rearrangements define a subtype [30-32]. In these

situations where the ground truth may be difficult to establish, computational pathology approaches for detecting molecular biomarkers from H&E-stained tissues may add yet another level of complexity. Thus, whilst the potential to implement molecular testing from digitised H&E-stained tissue is encouraging, significant validation is still required to determine whether computational pathology pipelines will demonstrate clinical utility.

Clinical guidelines recommend genetic testing in some patients with solid malignancies; unfortunately, these assays often have long turnaround times, are expensive, and have taxing sample requirements [13]. These limitations have so far hindered widespread genomic and molecular profiling implementation, particularly as the number of clinically actionable targets keeps increasing. There is a demand for inexpensive and scalable testing options that slot into the routine diagnostic pathway [13,23,33–36]. Since the genetic landscape impacts on a cancer phenotype, capturing the morphology of a lesion and its microenvironment by computational pathology offers a potential alternative [13]. Indeed, CNN models applied to routine H&Estained tumour sections have been described for the prediction of biomarkers such as mutations in KRAS [23,34], BRAF [33,34], TP53 [23,34,36,37], microsatellite instability [34], and tumour mutational burden (TMB) [35]. Moreover, weakly supervised CNN model-based frameworks utilise information from histopathology and other clinical reports as the ground truth label for an entire WSI in a classification or segmentation task [38]. Without a priori manual annotations, these networks learn to localise specific regions associated with a particular clinical, pathological, or genomic label [33,34,39,40]. One weakly supervised model discovered a new set of genotype-phenotype associations between histological tissue patterns and homologous recombination deficiency (HRD), a genomic aberration in a DNA damage repair pathway often found in cancers of patients with germline BRCA1/2 mutations. The AI model, trained to detect HRD, identified the enrichment of carcinomatous cells with clear cytoplasm and intratumoural fibrosis as a phenotypic signal of this deficiency [40].

The composition of the tumour microenvironment (TME) in solid cancers plays a key role in cancer initiation and progression [6]. Spatial analysis powered by AI has helped to decipher the TME and revealed pathways contributing to both immune escape and the lack of immune cell ingress [41]. Computational pathology methodologies capturing tumour-infiltrating lymphocytes (TILs) on H&E-stained breast and other cancers according to guidelines defined by the International Immuno-Oncology Biomarker Working Group, also called the TILs-WG [42], have shown potential in predicting clinical outcome [38,43–45]. The TILs-WG has organised a public grand challenge for computational assessment of TIL-counts alone and integrated into nomograms with established prognostic variables. In head and neck malignancies, AI-based methodologies

for detecting TILs have shown superiority in separating patients' outcomes compared with manual TILs scoring and a better delineation of stromal TILs from native lymphocytes in lymphoepithelial tissues such as the oropharynx [46]. Our group has repeatedly demonstrated that assessing immune responses in lymph nodes of triple-negative breast cancer (TNBC) patients adds prognostic value [43,47,48]. In particular, the formation of germinal centres and an expanded sinus surface area in a patient's lymph nodes are associated with longer intervals of disease recurrence [43]. By implementing a multi-scale CNN-based framework, germinal centres and sinuses on digitised H&E-stained axillary lymph node sections were robustly quantified, comparable with inter-pathologist assessments [48,49].

Computational pathology may offer opportunities to detect early signs of precancerous changes in normal tissue. The assessment of premalignant lesions and potential precursor lesions that may or may not become invasive in a patient's lifetime suffers from poor inter-observer reproducibility [50–54]. Grading of oral dysplasia [52,53] and ductal carcinoma *in situ* (DCIS) [38,51], a non-obligate precursor and risk factor of invasive breast cancer, are prime candidates for AI-based approaches. Given our limited knowledge of precancerous tissue characteristics, AI-assisted computational pathology pipelines may reveal new features in seemingly normal tissue and, as such, offer new tools for early cancer detection approaches.

Towards adoption in the diagnostic histopathological path

Whilst translational research in computational pathology has created a plethora of AI tools over the last decade, the clinical utility of these pipelines will ultimately be determined by the pathologists using them and their impact on improving patient outcomes. Similar to the application of AI-based analytics to mammography in breast radiology, there is an underlying question of whether AI-based diagnosis will be sufficiently accurate to be of value to histopathologists. The quantification of features that are time-consuming and poorly reproducible through manual assessment, such as the scoring of biomarkers like Ki67 or TILs (as described above) in breast or prostate cancer grading, is an area where pathologists are looking at computational pathology. For some of these features, image analysis is proven to be non-inferior to pathologists' interpretation, despite the recognised difficulties in standardising pre-analytical components and the added complexity of digital pathology (e.g. image variability due to using different scanners) [55,56].

Computational pathology's utility beyond diagnostics has so far largely been understudied. It can be readily incorporated into routine laboratory practice to strengthen existing quality control measures and quality improvement strategies. This is especially important in cancer diagnostics, where the quality of tissue staining is critical for accurate histopathological appraisal and precision grading of tumours. Computational pathology algorithms that first evaluate tissue sections for staining quality can thereby contribute to improved turnaround times by ensuring that histopathologists will receive slides that are optimal for reliable assessment. Additionally, they could potentially be used as a prescreening tool for enrolment of patients in clinical trials based on the morphology of their tumours. Automated AI systems may well recuperate some of a pathologist's time, but the ultimate benefit will be measured, similar to the clinical utility of genomic testing [57], by their likelihood of improving health care outcomes.

As well as application in the assessment of timeconsuming and relatively 'mundane' scoring tasks, computational pathology solutions may have value for 'triaging' cases [54,58]. For example, the detection of lymph node metastasis is critical for the diagnosis and staging of many solid tumours, and pathology modernisation programmes in hospitals have started to evaluate AI-based tools to support the pathologists' workload and improve the speed of assessment of microscopic examination [59]. Computational pathology pipelines need large-scale validation before obtaining regulatory approval; however, they will have a positive impact on cancer diagnosis turnaround times.

Current clinical pathways are based on the human interpretation of histopathology slides, and the role of AI algorithms in clinical practice needs to be ascertained. For example, invasive breast cancer is defined as HER2-positive if >10% of tumour cells show strong, complete membrane reactivity [60]. An AI algorithm could potentially distinguish between 8%, 10%, and 12% HER2-positive staining, something the human eye cannot discern. Since we do not know the clinical significance of these small deviations of HER2-positive staining, clinical trials must now be designed to validate AI-based HER2 scoring. Quality control (QC) and quality assurance (QA) programmes that are mandatory in histopathology services are also excellent examples for similar programmes that will be required with AI algorithms at their core. For example, programmes will need to consider periodic or continuous monitoring of the data being fed into an algorithm and will be essential to monitor the impact of both changes to the underlying data distribution (data drift) and the relationship between the input data and output variables (concept drift) on the model predictions and their deviations from expected outputs. The magnitude of these deviations and overall model drift are likely to guide periodic retraining of deployed AI algorithms so that they remain relevant for their respective application. While several of these concepts have long existed in the pathology laboratory to guarantee accurate assessment, guidelines ensuring that the AI algorithms retain clinical confidence must be implemented. The diagnostic quality model (DQM) was recently proposed to offer an operational framework to guide the implementation of computational pathology into clinical practice and to measure its nested impact at

the diagnostic test, procedure, laboratory, or healthcare ecosystem level [61].

Consistency of results across computational pathology tools on the same set of WSIs is a critical issue that will likely hinder their clinical utility and ultimate benefit to patients. The assessment of standard biomarkers, such as histological grade in breast cancer, has always suffered from moderate concordance across pathologists, yet is still applied in daily practices worldwide. Although a degree of discordance is generally accepted amongst pathologists due to the human nature of manual assessment, AI-based tools will not be given the same leniency, potentially due to several (un)conscious reasons (e.g. added costs for implementation and deployment). This is topped by a current issue, whereby different computational pathology solutions developed for the same task exhibit significant variability. One example is an AI-based tool for HER2-low detection [62], which as a consequence could lead to different centres treating a patient differently. The fragmented approach to standardisation and regulatory approval of different assays as companion diagnostics is of concern, as we have seen for the detection of biomarkers such as PD-L1 in TNBC, where different scoring methods, positivity thresholds, assay sensitivity, and multiple other factors lead to different positive prevalence rates. Using computational pathology tools as companion diagnostics in prospective clinical trials may be prone to divergent outcomes for selecting patients for treatment unless solutions such as concordance studies to harmonise similar AI-based companion assays from different manufacturers have been set in place [63]. Computational pathology tools must be subjected to similar evaluations before gaining regulatory approval, and reference materials need to be developed by healthcare authorities to facilitate robust analysis of the efficacy of these applications. Furthermore, formally determining the clinical utility of a prognostic or predictive biomarker currently requires prospective randomised trials. It is rather unlikely that trials will be designed to demonstrate that the AI-derived biomarker expression shows the same clinical utility as previously demonstrated. The question then arises: what is the best way to demonstrate the clinical validity of these tools? Recently, there have been studies showing that real-world data can give similar evidence to randomised clinical trials [64,65]. Observational studies, based on the collection of real-world data in our hospitals and biobanks, could potentially be leveraged to validate clinical utility for computational pathology tools.

A looming question remains as to whether clinicalgrade computational pathology tools need to be fully interpretable for medical diagnostics. AI-based models are commonly described as unaccountable 'black-boxes', due to the opaque nature of their decision process [1,8,38,66]. Explainable models require methods that abstract the exact underlying rules that form a neural network's decision. Such a requirement was challenged in a recent survey of 25 pathologists on their views towards digital pathology, which ultimately found that the majority doubted that this would hinder adoption [67]. As it seems, the algorithm predictions' robustness, validity, and consistency are likely more critical to pathologists for their routine clinical practice.

As AI progresses towards greater integration with diagnostic pathology, pathologists must become central members of the 'computational pathology' team. The successful adoption of AI algorithms in pathology is dependent on adequately training pathologists to gain knowledge and understanding of AI model training and familiarity with the relevant software. Incorporating AI subjects in clinical training and integrating AI models into existing postgraduate teaching platforms may help to overcome these challenges and, in parallel, entice the new generation of medical students towards computationally driven diagnostics. This will further encourage future histopathologists to embrace computational pathology as an adjunct diagnostic tool.

Challenges of building clinical-grade computational pathology systems

The road to clinical adoption of computational pathology is paved with a number of challenges, summarised in Table 1. Whilst a more detailed review can be found in the corresponding computational pathology article of this annual review issue by Professor Rajpoot's group [68], we highlight some key areas of concern focused on the pathologists' perspective.

Operationally, integrating computational tools into the current workflow of pathologists poses two significant challenges. Expensive infrastructure to digitise, store, and manage WSIs will need to be integrated into clinical pathways with minimal impact and disruption to laboratory daily practice [14]. Pathologists will require time and training to adapt to this new digital workflow. Technically, the need for large datasets to build scalable and robust AI and the lack of standardisation in data collection and processing hinder the development and validation of AI-based algorithms [39,66]. In particular, model generalisation is a challenge in AI and is compounded in pathology due to variations in the quality and staining of WSIs caused by the divergence in tissue preparation, staining protocols, and slide scanning machines used to digitise glass slides [1,8,14,66,69,70]. As such, technical variability must be represented in any training data, and some have argued a minimum of 10,000 WSIs is needed to build clinical-grade computational pathology algorithms [39]. While training on slides from multiple sources would provide a solid foundation for developing robust AI models able to perform across a wide range of real-world WSIs, sharing sensitive datasets across institutions and possibly country borders introduces logistical and legal challenges [66]. In the context of regulatory and ethical challenges, the use of patient data for research and development requires careful consideration and regulation to ensure privacy and consent. WSIs do not contain any identifiable

embedded patient data, but accompanying clinical and/or genomic information must be handled carefully [14,71,72]. Paramount to the successful adoption of computational pathology are standardised guidelines and regulations for integrating AI in pathology, including procedures on data acquisition, designing, training, and validation of models that consider ethical and patient privacy concerns [71,72]. Guidelines such as the 'Good Machine Learning Practice for Medical Device Development: Guiding Principles' from the Food and Drug Administration (FDA) in the USA [73] and the Medicines and Healthcare products Regulatory Agency (MHRA) in the UK have provided a generalised framework to build upon. The British Standards Institute (BSI) has recently developed the first BSI standard for AI in healthcare: BS 30440. However, currently, there is no specific regulatory pathway for the approval of AI software, with recent computational pathology device FDA approvals falling under the *de novo* pathway [27]. Ultimately, the reported ethical-legal concerns regarding the implementation of computational pathology in daily practice can be mitigated by establishing robust regulatory frameworks that specifically address issues pertaining to accountability and liability. As an adjunct diagnostic tool, however, computational pathology is still subject to the expert discretion of the histopathologist, who remains the ultimate custodian of diagnostic assessment.

Acquired datasets will, to some degree, represent underlying social biases and, in turn, feed those prejudices to the models [1,8,66]; this lack of diversity could perpetuate the underrepresentation of patients from minority groups and sub-populations with lower socioeconomic status [71]. With a lower exposure to histology from specific populations, models may be prone to more inaccurate diagnoses for these groups [66]. Therefore, creating fair and equitable systems is fundamental to providing the best care and building confidence and trust in computational pathology in healthcare.

Financially, the cost of implementing and maintaining computational pathology systems can be a barrier to adoption, especially in resource-limited settings. Even if there is a clear benefit to pathologists' efficiency and patient care, there needs to be a clear economic argument to motivate the relevant stakeholders that digital and computational pathology are beneficial. There are high upfront costs for the infrastructure. Although computational pathology can contribute to democratising care across countries with different resources, installing the digital infrastructure will be a barrier to using computational pathology in laboratories in middle- and lowincome countries. Simply put, we can use the blueprint from other recent advances (e.g. next-generation sequencing) that achieved economic sustainability by demonstrating clinical utility.

As we are still in the early adopter phase of computational pathology, its current value resides in AI-based research and education and to a lesser extent in its immediate clinical application. The next step in

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Challer categoi	ıge ry Challeng	e	Description and potential consequences of risk	Type of risk	Impact	Mitigation approach
Operati	onal Infrastructu	 Slow transition from conventional to a digital pathology workflow across all hospital pathology departments nationally 	Slow implementation of digital pathology hardware (e.g. WSI scanners and servers for storage) and software (WSI management systems)	Risk to patients	High	Establish long-term digital pathology economic benefits and secure sustainable funding programmes to pay for the upfront short-term and ongoing infrastructure costs
Operati	onal Laboratory technicia technical training	Inadequate training for laboratory technicians in new digital pathology protocols for WSI generation and management	If laboratory technicians in charge of slide preparation and WSI generation are not appropriately trained as part of the transition to new digital pathology workflows, quality of WSIs could be severely affected, which will affect the downstream efficacy and predictive power of deep learning models (see below Quality control)	Model risk/risk to patients	Medium	As part of the transition programme to digital pathology, teaching courses will be required to train technicians to work with the new digital pathology protocols
Operati	onal Pathologist technical training	Inadequate training of pathologists using computational pathology technology	Inadequate training for pathologists in computational pathology and deep learning concepts could lead to below-par quality of care and pathologist dissatisfaction. The latter could lead to fewer personnel entering the profession	Risk to computational pathology adoption/risk to patients	High	Integration of computational pathology and basic deep learning topics into existing teaching syllabuses
Technic	al Quality cont	rol Inadequate standardisation of tissue preparation and scanning protocols	Differences in preparation protocols, including tissue fixation, cutting, and staining procedures, across different laboratories and slide artefacts may increase the complexity of a model's learning task and cause low-quality performance for particular tasks or cause models to overfit to training data (see below Model generalisation). Bad performance could hinder the required level of accuracy needed for clinical adoption of these pipelines and low generalisation capability may lead to performance degradation in clinically approved models which could negatively impact patient quality of care	Model risk/risk to computational pathology adoption /risk to patients	H	Quality control and assurance programmes in histopathology will need to be designed with computational pathology principles at their core. Pre-analytical protocols and data management should be standardised. Local calibration of models at each local centre may also mitigate intrinsic variability in datasets
Technic	al Model generalis:	Computational pathology deep learning models are unable to generalise to new WSI datasets in the clinic, causing unexpected, inconsistent, and inaccurate predictions leading to poor quality of care	Deep learning methods are prone to overfitting on training data such that they do not perform well on unseen external datasets that they have not seen before. This is compounded in computational pathology due to technical variation in slide preparation such as tissue fixation, cutting, and staining procedures that vary between laboratories. This variability can cause significant differences in morphology and introduce noise into model training that ultimately reduces the predictive capability of models and leads to errors in clinical and diagnostic decisions	Model risk/risk to computational pathology adoption/risk to patients	ч Э	As per Quality control above, quality control and assurance programmes will be required to standardise the preparation of slides that will help to minimise variability across institutions. Diverse datasets should be used for training to capture the heterogeneity that exists in real clinical data. Methods such as stain augmentation and generative-based deep learning models can be used to create synthetic data that capture and introduce a diverse spectrum of stain heterogeneity into datasets. Regulatory guidelines on appropriate external validation of clinical-grade models will help to identify where models do not meet the minimum requirements for clinical applications
Technic	al Datasets	Access to only small datasets that do not capture heterogeneity across real clinical scenarios and are therefore not appropriate to train and develop robust deep learning models	Getting access to large datasets for a single malignancy and patient population can be difficult, particularly from multiple centres, which introduces legal and contractual requirements for the sharing of patient data. Large-scale, multi-centric datasets are essential to build robust generalisable, non-biased algorithms that mitigate overfitting and capture the wide array of	Model risk/risk to computational pathology adoption/risk to patients	High	More standardised public datasets, such as the CAMELYON16 challenge, across different cancers should be developed to encourage research and best practice development for training and validating different tools for the same task. The research community should be encouraged to publish their datasets after publication of their work. Federated
						(Continues)

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Access to an developm models ft decrease and diagr and computat si widespres icy Loss of patie

Challenge category	Challenge	Risk	Description and potential consequences of risk	Type of risk	Impact	Mitigation approach
Ethical	Model bias	Biased models with low predictive power for underrepresented patient populations	Without careful curation of training datasets, models can learn underlying societal biases. This could lead to models that are prone to inaccurate diagnoses for converal annitorinos	Model risk/risk to patients	High	Careful curation of training datasets that capture the appropriate underlying patient population based on the model application. Periodic testing and validation will be recurrent to ensure that models stay fit for ourmose
Financial	Funding	Failure to secure funding for a health care system-wide transition from conventional pathology to digital and computational pathology	If sustainable investment and funding are not available to transform pathology departments towards digital and computational pathology, the benefits of these innovations for both patients and pathologists will not be realised. Funding will be needed for the initial costs of implementation and infrastructure and ongoing costs related to upgrading of infrastructure, storage and management of data, and training of staff	Risk to computational pathology adoption/risk to patients	hội	A clear economic and value-based argument for the adoption of computational pathology must be established to encourage governments and private capital to fund the digital and computational pathology transformation. Countries that rely on health insurance and reimbursement systems will need to adopt their current frameworks to incentivise the adoption of using computational pathology. The pricing of computational pathology systems will need to be determined and will depend on the potential clinical utility of Al biomarkers and their impact on reducing pathologist workload,
Cultural	Pathologist and patient attitudes towards AI	Rejection of Al-based innovation for clinical and diagnostic applications by both clinicians and the wider public	Adoption of computational pathology can only be successful with the acceptance of its use by pathologists and patients. From the pathologists' perspective, this first requires the transition to assessment of tissue using digital pathology, followed by the integration of Al software into their everyday workflow. If pathologists and the public cannot be convinced of both the benefits and the safety of these innovations, then such technology will never make it to	Risk to computational pathology adoption/risk to patients	Medium	reducing overall costs of care, and their impact on improving patient outcomes Including computational pathology as part of the standard training curriculum so pathologists understand the benefits and risks associated with deep learning in pathology and demonstration of the clinical value will be the optimal way to get their 'buy-in' for this technology. The cultural shift relies on local qualifications and availability of new skills. For example, departmental or practice leadership could consider training in change management. At a patient
			the clinic			level, continued education about artificial intelligence at a societal level will be required

establishing true innovations will be to ensure financial sustainability. Prospective trials will have to demonstrate the utility of computational pathology; however, additional steps include policy development, negotiation, and contracting, which are necessary to bridge the last 'valley of death' in the translation, utility confirmation, and integration of computational pathology innovations. The successful integration of computational pathology paradigms into clinical practice relies heavily on securing sustainable funding, which requires an understanding of the economic implications and local cost containment strategies. Countries utilising reimbursement models will need to adapt these frameworks to cover AI-assistive technologies, otherwise there may be limited incentives to deploy computational pathology software for diagnosis [27,66]. Thus, while the scientific advancements in pathology are groundbreaking, the financial sustainability of these innovations will be crucial to their success. Ultimately, a robust local network of digital pathology centres will facilitate economies of scale by expanding the computational pathology community and, in turn, advancing the technology and the regulatory environment to promote innovation.

A cultural shift is required for the successful adoption of AI-based tools in pathology, with buy-in for AI-powered diagnostics from both clinicians and patients. Patients will need to be comfortable with a diagnostics decision being driven by AI-based software, and pathologists will need to be at ease working in tandem with new tools to make clinical decisions. Inertia to a change in direction of established practice is common with technology. To start with, a paradigm shift from traditional to digital pathology needs to happen in our hospitals before computational pathology can be integrated into a clinical setting. Whilst the digital transition is underway in some countries, many have yet to begin. Continued translational research and clinical evidence demonstrating the predictive power of computational pathology and its positive impact on patient care will ultimately be the catalyst for pathology departments worldwide to make the transition [12].

Towards implementation of computational pathology

The application of computational pathology systems at scale will initially depend on adopting digital and computational pathology workflows as well as efficient laboratory information management systems in the local hospital setting (Figure 2). Ubiquitous adoption of digital pathology with appropriate infrastructure will fuel the scaling of computational pathology, including curating multi-centric large-scale WSI datasets capturing tissue staining heterogeneity and tackling current obstacles of poor generalisation of models. Standardised public datasets have become the norm for developing and evaluating AI models in other fields, such as the ImageNet Large Scale Visual Recognition Challenge in computer vision [1,75,76]. As computational pathology scales, public large-scale datasets [48] may become more readily available and virtual biobanks comprising large multinational cohorts can be created to advance reliable prospective and retrospective tissue-based cancer research globally. Increased robustness, consistency, and validity of AI models will further fuel confidence in computational pathology.

Federated and swarm-based learning techniques present a scalable network-based solution to leverage data from multiple centres without compromising data protection requirements. In a federated approach, data remain local to each centre and feed training of a corresponding local model. Intermediate results from local models are then aggregated by a single global model, obviating the need for any sharing of data beyond local firewalls (Figure 2) [77–80]. These strategies are emerging as strong contenders for addressing many challenges in developing AI models and could pave the way to build robust, generalisable, secure, less-biased, and collaborative computational pathology software [66]. High-quality datasets will help to develop models addressing diseases with lower incidence [81]. One such approach, continual learning, addresses the problem of 'catastrophic forgetting', enabling models trained on common carcinomas to adapt to smaller datasets from less common diseases whilst maintaining a robust memory of prior cancer [81].

Finally, the transformer engine of generative large language neural networks has surpassed previous architectures in natural language processing tasks and has begun to usurp CNNs in computer vision [82]. So far, applications of these networks in computational pathology are in their infancy, but their power to model longdistance dependencies could unlock new insights into computational pathology [83,84]. At the time of writing, OpenAI released the latest in their GPT series, GPT-4, a multi-modal model that understands both text and images [85]. The interplay between natural language and vision of these powerful models could, although unlikely in the near term, lead to a fully automated, interpretable computational pathology solution that might learn from both clinical reports and WSIs to aid pathologists in their digital assessment and help to produce a full histopathology report.

The commercial landscape

In 2021, 5 years after the CAMELYON16 challenge, Paige (New York, USA) received the first FDA approval for an AI product in computational pathology for their Paige Prostate solution [8]. This year, the ARTICULATE PRO project will evaluate the deployment of Paige Prostate in the prostate cancer pathway at the Oxford University Hospitals (OUH) NHS Foundation Trust. IBEX Galen (Tel Aviv, Israel) [86] was recently awarded the Artificial Intelligence in Health and Care Award and will begin deploying their Galen Breast product in October 2023 to assess over 10,000 biopsies as part of routine practice across five



Figure 2. Towards a scalable network-based solution of utilising computational pathology. On the left, Centre 1 shows a schematic of a computational pathology workflow in a local setting. WSIs generated from daily practice, clinical trials, and registry trials are stored digitally in a cloud-based WSI repository. The pathologist can access computational pathology software, which sits on top of the AI analytical engine and the WSI management system and is connected to the local electronic health record system. External AI plugins can be added to the AI analytical engine using programmes such as Python, OpenCV, and QuPath. On the right, an exemplar of a federated learning framework to train a privacy-preserving multi-centric AI model using WSIs from three different centres is shown. Each centre trains a local model on its respective dataset and at the end of each iteration of training, intermediate results are sent to a global model for aggregation. The global model sends back the updated parameters to each local model for the next iteration of training. This framework respects data privacy as datasets remain at the local site and encourages the development of robust computational pathology systems trained on large multi-centric datasets. This figure makes use of the QuPath logo courtesy of the open source software QuPath [74] and the Python logo courtesy of the Python Software Foundation (https://www.python.org/psf-landing). This figure was created with the aid of BioRender.com.

NHS trusts in the Midlands (UK). Nevertheless, as of April 2023, the FDA had approved fewer than ten AIenabled medical devices in pathology, compared with nearly 400 in radiology. Similarly, only a small number have so far achieved Conformité Européenne (CE) marking in Europe. Besides Paige and IBEX, several computational pathology companies [e.g. PathAI (Boston, USA) [87], Aiforia (Helsinki, Finland) [88], Proscia (Philadelphia, USA) [89], Owkin (New York, USA) [90], and Panakeia (Cambridge, UK) [91]] are developing commercial products aiming to assist in both routine diagnostic tasks, ranging from breast and prostate cancer diagnosis, grading and subtyping, lymph node metastasis detection, to predicting molecular alterations, overall survival, and response to treatment. For instance, Aiforia has received CE approval for PD-L1 scoring, a biomarker of immunotherapy response [92]; Owkin has developed a model to predict the overall survival of malignant mesothelioma patients [93]; and

PathAI has developed a pipeline for the prediction of HRD from H&E-stained tissue [94]. The emergence of these nascent ventures illustrates the hope and opportunity of computational pathology and points to the areas within pathology where at least researchers and the private markets believe computational pathology can have the most impact. Globally, as pathology departments continue to 'go digital' in the upcoming years, the opportunities for computational pathology diagnostics will continuously increase. One of the main challenges with the imminent adoption of clinical computational pathology applications resides in the lack of appropriate regulatory approval pathways from agencies such as the FDA and the CE [27]. In the UK, the MHRA has announced the 'Software and AI as a Medical Device Change Programme' to ensure that regulatory requirements are fit for protecting patients and also encourage innovation by not burdening prospective ventures with high financial or regulatory costs.

Conclusions

AI-based tools have the potential to unleash a new era of precision medicine in oncological pathology. If the barriers to entry can be addressed, computational pathology will expand the influence of morphological assessment on diagnosis and treatment for cancer patients. All stakeholders will need to act, and system design must be led by pathologists as the ultimate users of these algorithms, whilst regulators, including pathology bodies, will need to produce guidelines and standards to harmonise best practices. Techniques such as distributed learning offer possibilities for building scalable AI models from large datasets without compromising data privacy. Over and above histology, combining computational pathology with molecular data from other nascent technologies (e.g. single-cell sequencing, spatial transcriptomics, and proteomics) will further advance our understanding of the mechanisms that govern disease and potentially inform on criteria for treatments. Finally, computational pathology systems, for at least the foreseeable future, should only be seen as tools, with ultimate medical decisions remaining firmly in the pathologists' hands. Utilising computational pathology and AI similar to advanced molecular technologies may eventually supplement traditional morphological approaches to improve patient care.

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Author contributions statement

GV and AG conceptualised the paper. GV, AG, JKL, SEP and RS prepared the main manuscript. DR, WN, ST, KPS, TN and SR provided content, detailed reviews and edits within their areas of expertise. GV and AG created the figures with support from JKL and RS. GV and AG provided overall supervision.

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