Precision Medicine, Artificial Intelligence, and Genomic Markers in Urology. Do we need to Tailor our Clinical Practice?

Medicina de precisión, inteligencia artificial y marcadores genómicos en urología. ¿Debemos cambiar nuestra práctica clínica?

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Abstract

Precision medicine plays a key role in urological oncology practice nowadays, with the breakthrough of the poly (ADP-ribose) polymerase inhibitors (PARPi), which play a critical role in different DNA damage repair (DDR) pathways, the immune checkpoint inhibitors, the genomic expression profiles and current genome manipulation-directed targeted therapy. Information and technology (IT) are set to change the way we assess and treat patients and should be reviewed and discussed. The aim of the present article is to demonstrate a detailed revision on precision medicine, including novel therapeutic targets, genomic markers, genomic stratification of urological patients, and the top-notch technological breakthroughs that could change our clinical practice.

We performed a review of the literature in four different databases (PubMed, Embase, Lilacs, and Scielo) on any information concerning prostate, bladder, kidney and urothelial cancer novel treatments with PARPi, immune checkpoint inhibitors (ICIs), targeted therapy with fibroblast growth factor receptor inhibitors (FGFRi), and theranostics with prostate-specific membrane antigen (PSMA) targeted monoclonal antibodies. Artificial intelligence, machine learning, and deep learning algorithm in urological practice were also part of the search. We included all articles written in English, published within the past 7 years, that discussed outstanding therapies and genomics in urological cancer and artificial intelligence applied to urology. Meanwhile, we excluded articles with lack of a clear methodology and written in any other language than English.

One-hundred and twenty-six articles of interest were found; of these, 65 articles that presented novel treatments of urological neoplasms, discussed precision medicine, genomic expression profiles and biomarkers in urology, and latest deep learning and machine learning algorithms as well as the use of artificial intelligence in urological...
practice were selected. A critical review of the literature is presented in the present article.

Urology is a constantly changing specialty with a wide range of therapeutic breakthroughs, a huge understanding of the genomic expression profiles for each urological cancer and a tendency to use cutting-edge technology to treat our patients. All of these major developments must be analyzed objectively, taking into account costs to the health systems, risks and benefits to the patients, and the legal background that comes with them. A critical analysis of these new technologies and pharmacological breakthroughs should be made before considering changing our clinical practice. Nowadays, research needs to be strengthened to help us improve results in assessing and treating our patients.

**Resumen**

La medicina de precisión juega un rol fundamental en la práctica clínica de la urología oncológica en la actualidad, con el desarrollo de los inhibidores de la poli (ADP-riboosa) polimerasa (PARPi), que juegan un papel fundamental en las distintas vías del reparo del ADN dañado (RAD), los inhibidores del punto de chequeo inmune (ICI), los perfiles de expresión genómicos, y la terapia blanco-dirigida a la manipulación genómica. El desarrollo tecnológico y la informática están cambiando la forma como evaluamos y tratamos a los pacientes, y se debe discutir y revisar a detalle. El objetivo de este artículo es hacer una revisión detallada acerca de la medicina de precisión, genómica, y los avances tecnológicos en nuestro campo.

Realizamos una revisión de la literatura en cuatro bases de datos diferentes (PubMed, Embase, Lilacs, y Scielo), buscando cualquier información relacionada con cáncer de próstata, vejiga, riñón y carcinoma urotelial, tratamientos novedosos con PARPi, ICI, terapia blanco-con inhibidores del receptor del factor de crecimiento de los fibroblastos (FGFRi) y teragnósticos con anticuerpos monoclonales dirigidos al antígeno de membrana específico de la próstata (AMEP). Inteligencia artificial, aprendizaje de máquinas y algoritmos de aprendizaje profundo en la práctica urológica también fueron revisados. Incluimos artículos escritos en inglés, publicados dentro de los últimos 7 años, que abordaran terapias novedosas y genómica en cáncer urológico e inteligencia artificial aplicada a la urología. Excluimos artículos con falta de una metodología adecuada y escritos en cualquier idioma diferente al inglés.

En total, 126 artículos de interés fueron encontrados, y, de estos seleccionamos 65 artículos que reportaban tratamientos novedosos para neoplasias urológicas, discutían medicina de precisión y perfiles de expresión genómica y bio-marcadores en urología, algoritmos de aprendizaje profundo, aprendizaje de máquina, y el uso de inteligencia artificial en la práctica urológica. Se hizo una revisión crítica de la literatura que se presenta en este artículo.

La urología es una especialidad constantemente en cambio, con un gran rango de avances terapéuticos, un gran conocimiento de los perfiles de expresión genómica para cada cáncer urológico, y una tendencia a utilizar tecnología de punta para estudiar y tratar a nuestros pacientes. Todos estos desarrollos se deben analizar objetivamente, y hay que tener en cuenta los costos al sistema de salud, los riesgos y beneficios para los pacientes, y el contexto legal que implica cada uno. Hasta la fecha, estos avances tecnológicos y farmacológicos se deben analizar con cautela antes de vemos en la posición de cambiar nuestra práctica clínica. Se debe fortalecer la investigación médica para mejorar los resultados en el tratamiento y abordaje de nuestros pacientes.

**Keywords**

- medicina de precisión
- genetica
- genomica
- inteligencia artificial
- teranosticos
- cancer
Introduction

New frontiers in urology are set further beyond with the new technological, genomics and pharmaceutical developments in this field. Urological practice has changed considerably in the last decade, new therapeutic targets have been studied, including manipulation of the immune system to attack cancer cells. The genomic expression profiles and genome manipulation-directed targeted therapy are constantly changing our practice and are set to be the base of a tailored approach for each patient’s genomic aberrations, also known as precision medicine.

Technology has had a major influence in urology in recent years, initially with the adoption of electronic records of the clinical history and with picture archiving and communication systems (PACS). Following with the development of deep learning algorithms, which are multilayered neural networks that learn from vast amounts of data, machine learning, which are algorithms fed by the exposition of data over time, with constantly improving performance and artificial intelligence (AI) which are computerized programs that can sense, reason, act and adapt according to an specific situation. All of these cutting-edge technological tools have been studied for the treatment of urolithiasis, urological cancer, hypospadias and have been able to successfully identify renal cell carcinoma, prostate carcinoma in surgical pathology and to discriminate tumors in white light cystoscopy.

The aim of the present article is to present a detailed revision on precision medicine, including novel therapeutic targets, genomic markers and genomic stratification of urological patients, and the top-notch technological breakthroughs that could change our clinical practice.

Methods

We performed a review of the literature in four different databases (PubMed, Embase, Lilacs, and Scielo) on any information concerning prostate, bladder, kidney and urothelial cancer novel treatments with poly (ADP-ribose) polymerase inhibitors (PARPi), immune checkpoint inhibitors (ICIs), targeted therapy with fibroblast growth factor receptor inhibitors (FGFRi), and theranostics with prostate specific membrane antigen (PSMA) targeted monoclonal antibodies. A search for prostatic adenocarcinoma (PCa) and urothelial carcinoma (UC) genomics was conducted. Artificial intelligence, machine learning, and deep learning algorithms in urological practice were also part of the search.

The search criteria were established in the form of free text and indexed terms. We used the MeSH terms: kidney cancer, prostate cancer, transitional cell carcinoma, poly ADP-ribose polymerase, antineoplastic agents, immunological, 177Lu-EB-PSMA-617, 68Ga-PSMA, intelligence, artificial, learning, deep, learning, machine, urologic diseases, urologic surgical procedures, urinary lithiasis, pathology surgical, genomics, comparative, genetic screening, and growth factors, fibroblast. The search was limited to publications in the past 7 years, and articles written in any other language but English were discarded.

A gray literature search was also performed on the pages of the National Technical Information Service (NTIS) and the European Association for Gray Literature Exploitation (EAGLE); however, no additional relevant information was found.

The articles were all original studies. References were reviewed by title and abstract by two independent reviewers. From the initial selection of articles, the references were reviewed integrally, ensuring they provided the aforementioned information of interest on all the topics. Duplicate studies were removed, and studies written in any language different from English were discarded.

Discussion

Precision Medicine and Genomic Markers

Precision medicine, despite being relatively new, is a concept that has been part of healthcare for decades. It takes into account the individual bases of genomics, lifestyle, and environment to precisely tailor personal therapeutic targets to treat disease and stratify patients to guide the best therapeutic approach.

Genome instability has been described as one of the hallmarks of urological cancer. In recent years, different DNA damage repair (DDR) pathways have been found to be altered in urothelial carcinoma (UC), renal cell carcinoma (RCC), and metastatic castration-resistant prostate cancer (mCRPC). New targets in urological oncology are emerging at a fast pace in the era of precision medicine.

Precision medicine is powered by patient data, health records, and genetic codes. Initiatives like The Cancer Genome Atlas research network (TCGA), which is a joint effort between the National Cancer Institute (NCI) and the National Human Genome Research Institute (NHGRI), have molecularly characterized over 20,000 primary cancers and generated over 2.5 petabytes of genomic, transcriptomic, proteomic, and epigenomic data, which has depicted the genomic landscape of non-metastatic and metastatic PCa.

Understanding of the genomic landscape in urological neoplasms has identified various dysregulated biological pathways that are relevant when determining the prognosis and natural course of the disease but that are very important for designing and using precise therapeutic targets to offer the patients a personalized management strategy according to their personal genomic alterations.

Four main pharmaceutical breakthroughs in urological cancer that are changing urological clinical practice worldwide are PARPi, ICIs, FGFRi, and theranostics with PSMA-targeted monoclonal antibodies with β and α-emitting radioisotopes.

Immune checkpoint inhibitors are designed to enhance or re activates antitumor immunity. It is one of the hot topics in urological cancer nowadays, given that they could be used for treating PCa, urothelial carcinoma, and renal-cell carcinoma (RCC), amongst others. Made easy ICIs could be divided in programmed death receptor ligand 1 (PD-L1) inhibitors, cytotoxic T-lymphocyte antigen 4 (CTLA-4) inhibitors and therapeutic vaccines such as sipuleucel-T, which are autologous mononuclear cells ex-vivo activated with a recombinant fusion protein fused to granulocyte-macrophage colony-stimulating factor.
factor, an activator of immune cells and PROSTVAC-VF ([Sanofi, Bridgewater, NJ]), a cancer vaccine made of engineered poxviral vaccine targeting prostate-specific antigen (PSA)-expressing cells, both in the mCPRC scenario.\textsuperscript{2,4,17} Ipilimumab is a monoclonal antibody that targets CTLA-4, a protein receptor for downregulation of the immune system. It has been studied for the treatment of metastatic renal cell carcinoma (mRCC) in conjunction with nivolumab (Checkmate 214), and it is currently under research in mCRPC.\textsuperscript{2,4,22} (\textbf{Fig. 1.})

Programmed death receptor ligand 1 (PD-L1) inhibitors are human monoclonal antibodies against programmed death receptor (PD-1) blocking PD-L1 from binding to PD-1 on activated T-cells, empowering the immune system to attack neoplastic cells.\textsuperscript{2,4,23} Some of the PD-L1 inhibitors are nivolumab, atezolizumab, avelumab, and pembrolizumab. Nivolumab has been studied in mRCC, showing improved overall survival (OS) compared with tyrosine kinase inhibitors (TKIs) (CheckMate 214, CheckMate 025).\textsuperscript{22,24} It has shown a meaningful clinical benefit in metastatic urothelial carcinoma, still phase 3 trials pending (CheckMate 275, CheckMate 032).\textsuperscript{25,26} but has shown no benefit yet in metastatic PCa.\textsuperscript{2,4} Avelumab has been recently reported to improve progression-free survival (PFS) in combination with axitinib versus TKI alone in mRCC (JAVELIN Renal 101).\textsuperscript{27} Atezolizumab has been widely studied in metastatic urothelial carcinoma (IMvigor 211) and mRCC (Immotion 151) with not very promising results.\textsuperscript{28,29}

Pembrolizumab is the only ICI approved by the Food and Drug Administration (FDA) for solid tumors, based on the presence of defects in DNA mismatch repair (MMR) genes leading to microsatellite instability (MSI), which is associated with a high mutational burden.\textsuperscript{2,4,23} It has been reported that 12\% of patients with advanced PCa have a hypermutated subtype, and it is correlated with MMR mutations (MSH2 or MSH6), which could also be found in up to 8\% of Gleason pattern 5 PCa.\textsuperscript{1,2,4,17} Pembrolizumab is currently recommended for mCRPC with MSI.\textsuperscript{2,4,23} In mRCC, it has been reported recently that pembrolizumab improves OS, and PFS in combination with axitinib (KEYNOTE-426).\textsuperscript{30} It has been extensively studied in metastatic UC as first-line therapy in cisplatin-ineligible patients (KEYNOTE-052)\textsuperscript{31} and second-line therapy after cisplatin-based chemotherapy (KEYNOTE-045), being associated with longer OS.\textsuperscript{32} Durvalumab, nivolumab, and avelumab had also been approved as monotherapy for advanced or metastatic UC after platinum-containing regimen failure.\textsuperscript{18}

Defects in DNA repair promote carcinogenesis through continued DNA replication without error correction and have been identified in up to 25\% of patients with mCRPC.\textsuperscript{1,2,33} The BRCA1 and BRCA2 genes are associated with higher Gleason scores, usually $\geq$ 8 and higher incidence of metastatic disease.\textsuperscript{1,2,17} Germline mutations in DDR genes, in metastatic prostate cancer, differ significantly from men with localized disease (11.8 vs 4.6\%). Specific mutations are BRCA2 (5.3\%), CHEK2 (1.9\%), ATM (1.6\%), BRCA1 (0.9\%), and PALB2 (0.4\%).\textsuperscript{1,17} (\textbf{Fig. 1.})

Poly (ADP-ribose) polymerase (PARP) play a significant role in DNA repair, they work by locating the DNA defect and binding to the replication fork until the repair begins.\textsuperscript{2,4,17,34,35} Poly (ADP-ribose) polymerase inhibition is one of the novel pharmaceutical targets in mCRPC and many other neoplasms given that through its inhibition the single-stranded DNA breaks will become double-stranded breaks that cannot be repaired and lead to apoptosis of the tumor cells.\textsuperscript{2,4,17,34–36} The potency of trapping PARP enzymes differ significantly between inhibitors, with a trapping efficiency following a downstream fashion: talazoparib, niraparib, olaparib, rucaparib, and veliparib.\textsuperscript{37} (\textbf{Figs. 2 and 3})

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{(A) Prostate cancer cell expressing programmed death receptor ligand 1 (PD-L1) inhibiting T-cell from attacking; cytotoxic T-lymphocyte antigen 4 (CTLA-4), a protein receptor for downregulation blocking T-cell through antigen-presenting cell interaction. (B) Anti-programmed death receptor and anti PD-1, blocking PD-L1 from binding to PD-1 on activated T cells, empowering the immune system to attack neoplastic cells; anti-CTLA-4, activating the T-cell to attack prostatic adenocarcinoma cells.}
\end{figure}
The last study published was the PROfound phase 3 open-label trial, which evaluated PARPi (olaparib) in men with mCRPC who had progressed receiving an androgen-signaling-targeted inhibitor (enzalutamide or abiraterone). All patients had at least one qualifying alteration with direct or indirect role with homologous recombination repair (HRR). One arm included gene alterations in BRCA1/2 or ATM. Radiological PFS was considerably longer in the PARPi group (7.4 vs 3.6 months) (hazard ratio [HR] 0.34; 95% confidence interval [CI], 0.25 to 0.47; \( p < 0.001 \)), and 81% of control patients who had progressed crossed over to the PARPi arm.\textsuperscript{35,38} The TRITON2 study of rucaparib in patients with DDR-deficient mCRPC showed a PSA and overall response rate of 53.6% and 47.5% respectively in BRCA patients.\textsuperscript{36,39} Thus, these findings encouraged the FDA to grant priority review of these two drugs.\textsuperscript{36,37,39}

The recently published TOPARP-B phase 2 trial (2019) aimed to evaluate the association between DDR gene alterations in mCRPC and response to olaparib 300 mg or 400 mg. Results showed that olaparib has antitumor activity against mCRPC with DDR alterations; the composite response was 54.3% in the 400 mg arm versus 39.1% in the 300 mg arm, suggesting that the ideal dose should be 400 mg, twice a day (at a expense of higher toxicity). It should not go without notice that all of these patients had already received docetaxel, 88 to 92%, a novel

![Fig. 2](image1)

**Fig. 2** (A) In human cells there are different DNA damage mechanisms. Single-strand breaks, these are repaired by the base excision repair pathway and double-strand breaks (DSBs), repaired by the homologous recombination repair (HRR) and the non-homologous end-joining pathways. BRCA ½ and ataxia telangiectasia mutated mutations directly affect the HRR pathway. (B) Poly-ADP-ribose polymerase (PARP) is a protein that carries surveillance within the cell and recognizes DNA damage, when identified, recruits HRR to repair DNA damage. Poly-ADP-ribose polymerase inhibitors bind to PARP1 and prevent identification of DNA damage, leading to DNA damage accumulation and cell death.
androgen-signaling–targeted inhibitor, and 31 to 45% cabazitaxel prior to the PARPi. There is a rationale that PARPi in combination with androgen-signaling–targeted inhibitor could improve PSA or radiologic response rate vs androgen-signaling–targeted inhibitor alone based on the premise that the androgen receptor (AR) pathways drive progression to mCRPC, which is due mainly to adaptive mechanisms that enable persistent AR signaling. Data shows that AR regulates transcription of several sets of MMR genes, PARP-1 can act as an AR co-factor and are upregulated in exposure to androgen deprivation therapy (ADT).

Stratification and molecular subtype identification in UC have gained popularity in recent years. The TCGA and several studies have shown that gene-expression datasets of UC suggest it could be classified in 6 molecular subtypes: NEURAL, LUMINAL, papillary-like (PAP), HER2L, mesenchymal-like (MES), squamous-cell carcinoma (SCC) and bladder carcinoma subtype of large meta-cohort databases (BOLD). Regarding muscle-invasive bladder cancer (MIBC), Choi et al identified two intrinsic subtypes: one basal and two luminal. The luminal I subtype has been associated with poor response to ICI, given its low gene-expression associated with immune response and a poor expression of PD-L1. Fibroblast growth factor receptor is involved in network signaling that regulate the cell cycle through proliferation, migration and differentiation processes. Luminal MIBCs are enriched with high levels of FGFR3 and activating FGFR3 mutations. Erdafitinib, a potent TKI of FGFR1–4, was evaluated in a phase 2 clinical trial in patients with locally advanced or metastatic UC with prespecified FGFR3 or FGFR 2/3 fusion mutation, who had received at least one cycle of systemic chemotherapy or were ineligible to cisplatin. They reported a response rate of 40%, 37% with a partial response and with a median OS of 13.8 months and median PFS of 5.5 months, which is higher than with taxanes or vinflunine and ICIs.

Theranostics in PCa is another field of interest in current urological practice and are poised to transform the treatment of patients with mCRPC. In simple words, theranostics is the combination of a targeted therapeutic agent with diagnostic tests such as ⁶⁸Ga-PSMA-11 PET/CT. VISION trial is a phase 3 multicenter prospective trial enrolling patients with progressive mCRPC with a positive ⁶⁸Ga-PSMA-11 PET/CT, treated with at least one novel ASI and one prior taxane-based regimen. The patients are randomized to ¹⁷⁷Lu-PSMA-617, a β-emitting radioisotope that releases β-particles that travel less than 2 mm and can irradiate small tumors in metastatic sites; at the same time, it emits low-energy gamma particles that are of utility while doing imaging-based tumor localization. Investigators planned to complete data collection in May 2020. Alpha particles targeting PSMA are in development and are a promising option for theranostics in the future, and currently a phase 1 multicenter study of the α particle emitter thorium-227 conjugated to a PSMA targeted monoclonal antibody is enrolling patients (►Fig. 4)

**Artificial Intelligence, Machine learning and Deep learning algorithms**

Improvements in prediction tools for disease behavior and treatment response are a recent area of interest in urology.
Artificial intelligence systems are setting new horizons in urological practice.

A machine learns when it changes its structure in response to external information based on algorithms, which ultimately means it will improve future performance and can be applied to any situation in which repetitive data can be obtained. Artificial intelligence is the application of machine learning and uses complex mathematical models to generate conclusions. Deep learning is a subset of machine learning, based on the human neuronal structure, that can generate data driven models of biological systems.\(^5,8\)

There are several AI methods that provide decision support systems (DSS), including Bayesian networks, expert systems, artificial neural networks (ANN), modeling systems, and decision trees.

Machine learning includes models that require the input of frequency data and previous knowledge or expert opinions, such as the Bayesian networks that allow to combine this information to arrive to an intelligent solution and expert systems that analyze information according to a series of rules or questions provided by an expert and generate conclusions. However, it cannot accommodate new questions or data.\(^5,8\)

Deep learning uses a multi-layered structure of algorithms that correspond to artificial neurons that run in parallel and can reorganize complex patterns according to a weight that is assigned to each input and obtain “knowledge” by feeding the information repeatedly to the system (ANNs and NFMs).\(^5,8\)

The mechanisms of AI are being widely studied and developed for improving diagnostic accuracy and prediction of disease behavior in urological conditions.

Regarding prostate multiparametric magnetic resonance imaging (mpMRI), AI algorithms are focused on the automated detection of suspicious regions and are expected to reduce reader interpretation times, increase performance of non-expert radiologists, and, ultimately, increase sensitivity and reduce inter-reader variability.\(^7\)

One study found the sensitivities of mpMRI-alone and computer aided diagnosis (CAD), based on a traditional machine-learning algorithm, were similar (79% vs 76%), but the greatest benefit of CAD was found to be in transitional zone
(TZ) lesions and for moderately experienced readers (84% vs mpmMRI-alone 67%, $p = 0.055$). Times of reading improved in CAD (4.6 vs 3.4 minutes, $p < 0.001$), and for PI-RADS V2 of 3 or more, sensitivity was superior in the CAD group than in mpmMRI-alone (72% vs 45%, $p < 0.001$).

Deep learning algorithms have been less studied for mpmMRI; however, some have proposed segmentation of the sequences to improve detection rates, assigning two parallel networks for T2 and ADC specifically, which has demonstrated an improvement in sensitivities compared with previous studies proposed. These studies are very promising; however, verifying the results of mpmMRI with the actual pathology is the real challenge. Studies have shown that mpmMRI signal characteristics are associated with tissue composition and density, specially the glandular components, which allows the creation of "radiopathic maps to distinguish cancerous regions.""

Segmentation of the histopathological components (stroma, nuclei, epithelium, lumen, etc.), which is known as semantic segmentation, is the basis of machine learning for pathology specimens. This allows to generate dichotomous results (benign vs malign) and further classify the tissue according to its characteristics generating a Gleason score. Based on these mechanisms described, deep learning systems (DLSs) have been developed to improve Gleason scoring, as follows. A retrospective study evaluated a DLS to grade prostate biopsies following the Gleason grading standard. It was designed to delineate individual glands and assign a Gleason pattern, grade, and group. This system was validated with 550 biopsies and compared with the results of 13 pathologists and 2 pathologists in training. It was tested in 579 biopsies and achieved a high correlation with the reference standard (Cohen kappa 0.918), differentiation between malignant or benign, Gleason grade, group 2 or 3, and it even outperformed 10 of 15 pathologist observers.

Another DLS was developed using 112 million pathologist annotated image patches and validated on 331 slides, compared with pathologist experts. It demonstrated a significantly higher accuracy of 0.7 ($p = 0.002$) and trended to a better risk patient stratification.

In recent years, DLSs, specially CNNs, have improved accuracy in image recognition, object detection, and semantic segmentation in kidney cancer diagnosis and classification. One study evaluated a CNN for RCC classification and survival prediction based on The Cancer Genome Atlas (TCGA) project that has resulted in digital haematoxylin and eosin (H&E) whole-slide images (WSI) of RCC. The CNN managed to classify each RCC from normal to three subtypes of RCC (AUC 0.93), which ultimately translates in survival prediction outcomes.

Furthermore, a paper proposed a DLS for the classification of kidney cancer into subtypes using the genome of 25 types of miRNA identified to determine tumor characteristics, information provided by the TCGA. A recurrent neural network is used to classify a miRNA sample into five cancer subtypes, with an accuracy around 95% and correlation coefficient of 0.92.

Regarding bladder cancer, the identification of the tumor in cystoscopy is the key for optimal transurethral resection of bladder tumor (TURBT). A study aimed to improve tumor localization and surgical resection in cystoscopy studies. With this purpose, videos with histologically confirmed tumors were selected and manually inserted into a platform based on convolutional neural networks constructed (CystoNet). Ninety-five patients were used for algorithm training, 5 for testing, and then it was validated in 54 patients. The sensitivity and specificity per-frame of CystoNet was 90.9% and 98.6%, respectively (detected 39 of 41 papillary and 3 of 3 flat bladder cancers).

Another DLS aims to predict the survival according to bladder cancer subtypes, using TCGA dataset of mRNA, miRNA, and methylation to infer two survival subtypes and apply it to any new individual sample. The high-risk survival subgroup had KRT6/14 overexpression and PI3K-Akt pathways.

Urinary stone disease is a highly prevalent condition, and the analysis of the size and volume of stones in the kidneys is an important point in surgical decisions and planning. Three-dimensional stone segmentation software, out of non-enhanced computed tomography (CT) have been compared with radiologists readings and promise more accurate results. Also, approaches to identifying the composition of stones have been described, with the purpose of better metabolic management of urinary stones, and an accuracy of 100% was achieved in differentiating uric acid from non-uric acid stones in a study that compared different AI algorithms for CT scan interpretation.

A review about the role of AI in urinary stone disease proposes these algorithms as the future of urology. Different methods are described, proposed to predict the probability of urinary stone disease out of symptoms, enhance the urinary stone tracking during shock wave lithotripsy (SWL) and minimize the number of emitted shockwaves during this procedure and also others that predict treatment success.

Several ANNs have been designed to predict the outcomes after surgical interventions as SWL and percutaneous nephrolithotomy (PCNL). One of them used the information on 454 patients (200 for training set and 254 for test set), to assess the relevance of clinical preoperative parameters on postoperative results (PCNL) by comparing them to the actual (observed) outcomes; the accuracy and sensitivity of the system was found to range between 81% and 98.2%, and it was able to predict stone-free rates with an accuracy of 86%. Stone-free rates after SWL were also evaluated by an ANN system based on information of 139 patients that was able to predict this outcome with an accuracy of 88.7%.

Finally, it is due to highlight the role of AI in surgical training. It has been studied mainly in robotic and laparoscopic surgery, with emphasis in anatomical landmark recognition as a fundamental step in automated surgery. Color and texture evaluation has been assessed during prostatectomy aiming to identify basic anatomical landmarks.

Also, the identification of instruments and detection of its movement promises great results regarding prediction of surgical skill and technique. Khurshid et al reviewed the videos of 12 surgeons anastomoses (robotic radical prostatectomy) using the global evaluative assessment of robotic skills (GEARS) tool, initially manually, by 25 peer surgeons and compared the scores with the results of a linear support
vector machine (SVM), which achieved an accuracy in classification of surgical skill of 83.3% and improved to 91.7% when joint movement was assessed. Finally, when the contralateral instrument was evaluated, accuracy was 100%.

**Conclusions**

Urology is a constantly changing specialty. Each urologist has the responsibility to update him- or herself with the novel breakthroughs in the field, be aware of new pharmaceutical developments, and, in this century, they need not only to be aware, but be involved in the cutting-edge engineer and technological advances related to their specialty. All of the aforementioned pharmaceutical targets, genomic markers and technological innovations must be analyzed objectively, taking into account costs to the health systems, risks and benefits to the patients, and the legal background that comes with them. Technological advances, such as artificial intelligence and deep learning algorithms, have shown promising results but could not replace physicians’ perspectives at the time. Certainly, our clinical practice needs to be retailed, and, with time, precision medicine is going to become the only possible medicine, tailoring treatments according to the genomic expression profile of each patient. Nowadays, research needs to be strengthened to make us reconsider changing the way we assess and treat our patients.

**Conflict of Interests**

The authors have no conflict of interests to declare.

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