Review Article

Systemic treatment of penile squamous cell carcinoma—hurdles and hopes of preclinical models and clinical regimens: a narrative review

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Abstract: Despite contemporary research efforts, the prognosis of penile squamous cell carcinoma (PeSCC) has not significantly improved over the past decade. Despite frequently encountered patient-related delayed medical consultations impairing outcomes, several other aspects contribute to the lack of advancement in the treatment of this condition. One essential reason is that translational research, a prerequisite for the clinically successful disease management, is still at an early stage in PeSCC as compared to many other malignancies. Preclinical experimental models are indispensable for the evaluation of tumor biology and identification of genomic alterations. However, since neither commercial PeSCC cell lines are available nor xenograft models sustainably established, such analyses are challenging in this field of research. In addition, systemic therapies are less effective and toxic without decisive breakthroughs over recent years. Current systemic management of PeSCC is based on protocols that have been investigated in small series of only up to 30 patients. Thus, there is an unmet medical need for new approaches necessitating research efforts to develop more efficacious systemic strategies. This review aims to highlight the current state of knowledge in the molecular alterations involved in the etiology and ensuing steps for cancer progression, existing preclinical models of translational research, clinically relevant systemic protocols, and ongoing clinical trials.

Keywords: Penile squamous cell carcinoma (PeSCC); treatment, biomarkers; copy number alterations (CNAs); mutational profiling

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Introduction

In contrast to the developing world with the incidence of 2.8–6.8 per 100,000 men, penile squamous cell carcinoma (PeSCC) is an uncommon tumor entity in Western countries where the incidence can be as low as 0.3 per 100,000 males (1). A plethora of risk factors have been identified including lack of neonatal circumcision, phimosis, chronic inflammation, lichen sclerosus, socioeconomic status, obesity, smoking, psoralen UV-A phototherapy, premalignant lesions as well as impaired immune response (1,2). Moreover, human papilloma virus (HPV) has been attributed to nearly 40–50% of cases (3,4). The disease most commonly affects men in their 6th decade of life, however young males may be at a higher risk for more aggressive tumor characteristics if diseased (5).

Dauntingly, contemporary reports from the USA, France, and Norway have shown that survival of PeSCC patients remained unchanged over respective periods of 10, 20, and 60 years despite all efforts in the amelioration of diagnostic and therapeutic paradigms (6-8). It is furthermore noteworthy that most currently used cytotoxic concepts for this less chemoresponsive condition have been introduced 10 years ago. Besides frequently encountered delayed medical consultations impairing oncologic outcomes, several other aspects contribute to the lack of progress in the treatment of PeSCC (9). One of the reasons is that translational research, a prerequisite for the clinically successful disease management, is still at an early stage in PeSCC as compared to many other malignancies. Indeed, fundraising for PeSCC preclinical research is challenging and often only possible when applying for funding solely dedicated to rare cancers hampering clinical advances as well. In the western world, this orphan disease is not “appealing” for drug development by the pharmaceutical industry due to its rarity (10). Of note, only 300 orphan drugs and devices were approved in the last 25 years in the US being literally only a drop in the bucket compared with the many thousands of orphan diseases (11).

Besides causing a lack of awareness, the rarity of the disease impedes an adequate conduct of translational research. It almost seems impossible to gather enough patients ensuring pertinent tissue biobanking and providing studies with sufficient statistical power, both compulsory requirements for identification and approval of new therapeutic regimes.

In this review we sought to shed light on the current knowledge of molecular alterations involved in this disease and potential molecular markers, preclinical research advances and available evidence for clinically relevant systemic treatment strategies in PeSCC. We present the following article in accordance with the NARRATIVE REVIEW reporting checklist (available at http://dx.doi.org/10.21037/tau-20-945).

Molecular profiling in PeSCC

Cancer genome is driven by the accumulation of somatic and structural variants, copy number alterations (CNAs), deregulation of gene expression, and epigenetic changes. Molecular classification of several tumor types has resulted in clinical applications and targeted therapy with the development of a new area named precision medicine. Although molecular classification is a reality for many tumor types, such as breast cancer, the scenario of precision medicine in PeSCC is still in its infancy.

Molecular profile of PeSCC has been recently reported using high throughput strategies to characterize the disease. However, this knowledge is limited for a few genetic and epigenetic studies resulting in potential prognostic markers with limited application in the clinical practice (Figure I). The main issues are the number of cases studied, the absence of data confirmation in large cohorts of tumors and functional models validating the findings, which could bring important surrogate markers of prognosis or be used as novel therapeutic targets.

Recurrent CNAs

CNAs are somatic-acquired changes of chromosome structure, leading to gains or losses of the genomic segment. These changes in gene dosage are frequently detected in cancer, with recurrent alterations associated with particular cancer types. CNAs occur in a large fraction of the cancer genome, activating oncogenes and inactivating tumor suppressor genes (12,13). However, CNAs were also mapped in regions without known genes associated with cell proliferation. In a pan-cancer study, Zack et al. described amplified regions without known oncogenes that were in turn enriched for genes involved in epigenetic regulation (14). Moreover, several CNAs are mapped close to metabolic cancer genes (15), which has significant clinical relevance in response to cancer treatment. In an analysis of 17,879 tumors from patients with known outcomes, Smith and Sheltzer detected 108 significant associations between gene CNA and outcome, compared to 23 associations between mutation and outcome (16). Among 28 of 30
Copy number alterations

Genome-wide copy number variation analysis

PPARG, LAMP3, TNFSF10, DCL1, MYC, EGFR, ERBB3, ART, PIK3CA, PTEN, COX2

DNA methylation

Cytosine Methyl Cytosine

Bisulfite conversion

PCR amplification

Pyrosequencing analysis

CDKN2A, RASSF1, DAPK, MGMT, RARB, RUNX3, TWIST1, RSOP2, SOX3, SOX17, PROM1, OTX2, HOXA3, MEIS1, BNGE

Pathways analysis

Genome-wide gene expression analysis (network analysis)

Pathways:
P13K/AKT/mTOR
RTK/RAS/MAP-kinase
Wnt/β-catenin
matrix-metalloproteinases

Figure 1 Schematic representation of different technological approaches to study genetic and epigenetic alterations in penile cancer (A) gene dosage analysis and detection of copy number alterations, (B) mutational profiling, (C) DNA methylation changes, and (D) transcriptomic and integrated pathways analysis. Representative examples of genes involved in the development and progression of PeSCC are indicated in rectangles. PeSCC, penile squamous cell carcinoma.
cancer driver genes, CNA was a significant prognostic factor.

The first report on DNA CNAs via array-CGH (comparative genomic hybridization) in PeSCC (N=38) described frequent losses of 3p and 8q, which were associated with advanced T and clinical stage, recurrence, and death from the disease (17). CNAs of genes mapped at chromosomes 3 (PPARG, LAMP3, and TNFSF10) and 8 (DLC1) were translated into altered expression levels and related to a worse prognosis. Gains of 8q24 detected by array-CGH included the MYC oncogene (8q24), which was confirmed as altered using fluorescence in situ hybridization. For the first time, the authors described different genomic profiles based on the presence or absence of the HPV infection, suggesting distinct etiologies for this disease. A subsequent study evaluated 64 PeSCC and confirmed differences in their CNA profile (recurrent gains of 1p, 3p, 5p, 8p and losses of 2q, 3p, 11q) between HPV positive and negative cases as well as based on tumor grade and lymph node status (18).

Recently, Macedo et al. performed array-CGH in 20 of 55 PeSCC (81.8% of them were HPV positive) (19). Despite of not having any significant clinical-histopathological association, a noteworthy number of altered chromosomal regions coincided with sites of HPV integration into the human genome. Gains of ERBB3 and EGFR, and losses of AKT2, PIK3CA, and PTEN were frequently described (>50% of cases). Increased expression of EGFR, COX2, PGE2 proteins, and decreased p53 expression were reported among 43 PeSCC tested by immunohistochemistry. According to these authors, EGFR and ERBB3 amplifications are potentially involved in the development or progression of PeSCC positive for high-risk HPV.

Overall, these studies showed significant differences in the genomic profiles of PeSCC according to HPV infection status, suggesting that these tumors probably require different treatment strategies. These initial studies showed distinct genomic profiles of HPV-positive and -negative PeSCC. Moreover, different potential prognostic markers were investigated in only a small number of cases. Taken together, the role of CNAs in PeSCC still needs to be better explored.

Mutational profiling

Advances in sequencing technologies have been instrumental in identifying new targets with potential therapeutic value in various tumor types. In PeSCC, several studies have reported somatic mutations in TP53, CDKN2A, PIK3CA, HRAS, KRAS, CCND1, and STK11 (20-27). Although in a small cohort of PeSCC, these studies revealed that EGFR amplification and CSN1 mutations were detected in primary tumors and lymph node metastasis (22,24,25). Ferrándiz-Pulido et al. evaluated somatic mutations in genes downstream of EGF receptor in 10 in situ and 65 invasive PeSCC (23). They reported frequent PIK3CA, HRAS, and KRAS mutations in p53-negative tumors, suggesting a role of mTOR signaling activation in PeSCC. An analysis of 67 PeSCC, phosphorylated mTOR overexpression was associated with lymph node-positive and HPV-negative tumors. Further studies described the role of PI3K/AKT/mTOR giving support for strategies of treatment using inhibitors of this pathway in PeSCC (28,29).

Using protein expression in a tissue microarray composed of 57 cases, Azizi et al. reported that PI3K-AKTmTOR pathway up-regulation and HPV positivity were associated with a favorable prognosis of PeSCC patients (29). These preliminary data suggested that mTOR pathway proteins have the potential to stratify PeSCC patients, which might help in treatment decisions.

Whole exome sequencing was recently reported in 30 PeSCC matched with normal blood samples (27). A total of 827 mutated genes was found; 94 of them presented recurrent variants, including FAT1, HRAS, NOTCH1, and CASP8 (found in 4 cases each). Other mutations were described in genes frequently reported as altered in PeSCC, such as TP53 and PIK3CA. The RTK/RAS/MAP-Kinase pathway was frequently altered (46% of cases showing gene variants). In this cohort of cases two pathways involved in inflammatory response (Notch in 30% of altered cases and Hyppo in 23.3%) were also altered (27). Formalin-fixed paraffin-embedded metastatic PeSCC (N=78 cases) showed frequent alterations in mTOR (NF1 and PTEN), DNA repair (ATM and BRC42), and tyrosine kinase (EGFR, FGFR3, and ERBB2) pathways (26).

These results give additional support for target-specific therapies. For example, patients having PeSCC with NOTCH1 loss of function could be treated with PI3K/mTOR inhibition. The involvement of inflammatory pathways indicated treatments based on immunotherapy combined with chemotherapy. Patients whose PeSCC present alterations in genes involved in DNA repair (such as ATM and BRC42) and tyrosine kinase (ERBB2, FGFR3, and EGFR) pathways could also benefit from targeted therapies. Application of high throughput techniques renders a massive
amount of data, providing unprecedented opportunities for identifying predictive markers of therapy response or therapies based on specific targets. Molecular studies in PeSCC are restricted to a limited number of cases and there are no comprehensive studies involving multiple platforms of the same case, which are critical to the establishment of targeted therapy. However, the results obtained to date are encouraging, leading support that patients could potentially benefit from target-specific treatments.

**DNA methylation**

Nowadays, it is recognized that epigenetic changes are critical during cancer development and progression. Epigenetic modifications, including DNA methylation and covalent histone modifications, are commonly disrupted in cancer cells. The best-known and studied epigenetic marker is DNA methylation. In the human genome, the cytosine methylation occurs almost exclusively at CpG dinucleotides across the genome in different contexts (CpG islands or non-CpG islands such as shore, shelf, and open sea regions). Both, global hypomethylation and focal hypermethylation at promoter-associated CpG islands are commonly observed in cancer cells. Moreover, DNA methylation participates in complex chromatin interactions networks and can modify gene expression. Aberrant DNA methylation has a critical role in tumorigenesis and has been reported in all tumor types. Changes in DNA methylation have been associated with resistance to cancer therapy (30). DNA methylation-based biomarkers are proposed as useful markers to stratify cancer patients according to prognosis (31). The methylation pattern of specific genes, including CDKN2A, RASSF1, DAPK, MGMT, RARB, RUNX3, has been reported since 2003 [for review (32)]. To date, DNA methylation using genome wide analysis was described by Kuasne et al. (32) and Feber et al. (33). The DNA methylation profiling in 38 PeSCC matched with 11 surrounding normal tissues revealed a hypomethylated profile in tumors without lymphatic involvement (33). These authors also reported an epigenetic HPV signature able to predict the HPV status and survival in independent cohorts of head and neck and cervical cancers. The second study describing genome wide DNA methylation analysis used the approach of data integration between DNA methylation data and transcriptome results (25 PeSCC, 10 surrounding normal tissues, and 4 normal glans) (34). The authors found a panel of 54 genes (such as TWIST1, RSOP2, SOX3, SOX17, PROM1, OTX2, HOXA3, and MEISI) with an inverse correlation between DNA methylation and gene expression. These findings pointed out that DNA methylation drove the regulation of pathways associated with the penile carcinogenesis, including embryonic stem cells, cell cycle, immune response, and Wnt/β-catenin signaling. Cytokines (such as IL1A, IL1B, TNF, CXCL1, CCL20) and MMPs (MMP7, 9, 10, 12, 13) were upregulated in PeSCC without correlation with HPV positivity. The authors described that BNDE hypomethylation was associated with lymph node metastasis and shorter disease-free survival (33). Although the number of cases was limited and the platform interrogated 27K CpGs, distinct methylome and transcriptome profiles were found according to the HPV status, suggesting that distinct therapeutic strategies should be applied to these patients.

**Transcriptomic and integrative analysis**

Studies on gene expression analyses in PeSCC are scarce. Kroon et al. evaluated 56 PeSCC according to lymph node metastases (36 positives and 24 negatives) using a 35K gene expression platform (35). A 44-probe classifier correctly grouped 29 of 30 cases (96%) in the training set, while for the validation group, only 14 of 26 samples (56%) were correctly classified. Expression array findings were also reported in data integration studies, as described above by Kuasne et al. (2015). Two other studies integrated transcriptomic data with miRNA, CNAs, and DNA methylation results (36,37). Kuasne et al. (37) performed an integrative analysis in 23 PeSCC and 12 non-neoplastic tissues using miRNAs (TaqMan Human MicroRNA Assay System Set v.2.0, Applied Biosystems) and mRNAs (Whole Human Genome 4x44K, Agilent Technologies) expression data. From the gene lists generated in these analyses, the authors selected 8 miRNAs and 10 transcripts for validation using RT-qPCR (array independent samples: 36 PeSCC, 20 surrounding normal tissues, and 10 normal glans). MMP1, which is supposed to be regulated by hsa-miR-145-5p, was described as a predictive marker of lymph node metastasis. Moreover, PPARG, VEGF, EGFR, and matrix metalloproteinase pathways were dysregulated, highlighting their involvement in PeSCC. A multidimensional integrative analysis (CNAs, DNA methylation, miRNA, and mRNA expression) was described by Marchi et al. in 20 usual PeSCC (36). Ten top genes among 16 driver candidates (AR, BIRC5, DNMT3B, ERBB4, FGFR1, PML, PPARG, RB1, TNFSF10, and STAT1) showed deregulation in a validation set of 33 PeSCC samples. **BIRC5** and **DNMT3B** up-regulation was associated with a shorter overall survival.
The current knowledge of the transcriptomic analysis in PeSCC remains limited.

Models for preclinical research

Tumor cell lines, patient-derived tumor xenografts, and genetically engineered mouse models are meaningful approaches to evaluate the efficacy of potential therapies as well as the mechanisms of treatment resistance. Cell lines have been extensively used in cancer research and, although they have significantly contributed to the cancer research advancements, several limitations and challenges are known. Establishment of cell lines is time-consuming, the architecture of the primary tumors is not reproduced, contamination of fibroblasts is frequently observed, and the protocol used to eliminate them is laborious. Furthermore, in many cases, in vitro cell selection results in significant differences at molecular levels comparing the primary tumor and its derived cell culture. To overcome these challenges found in two-dimensional cell lines, xenografts and three-dimensional cell cultures recapitulate components of the tumor environment. In xenograft models, tumor cells or tumor itself are transplanted into immunodeficient animals. The study in xenografts requires significant expertise to maintain animals and perform experiments. Though the number of animals in research is limited, protocols are expensive and not suitable to be used in large-scale drug screening assays (38,39).

Tumor cell-derived cultures in 3D (also named organoids or tumoroids) are valuable models for predicting therapeutic response (40,41). The 3D-cell culture is based on the use of Matrigel, a gelatinous protein mixture secreted by Engelbreth–Holm–Swarm mouse sarcoma cells that resemble complex extracellular environment (42). Alternatives for the use of Matrigel have been proposed as a scaffold-free model using microcavity technology. Recent studies have demonstrated that tumor organoids (tumoroids) recapitulate histology, gene expression, and genomic profile of the original tumor (40,43-45). Screening of drug sensitivity in established tumoroids demonstrated their potential for implementation in clinical practice as a guide for individualized medicine (46,47). Tumoroids are usually established from resected tissue of the primary tumor, and consequently, one of the main challenges is to capture the substantial heterogeneity of tumors in vivo. However, tumor heterogeneity could be studied in tumoroids clonally established from single cells derived from tumor tissue (48).

Few primary PeSCC cell lines have been reported, with none being commercially available. Naumann et al. performed 2D-cell cultures from nine primary tumors, three lymph node metastases and one distant metastasis from 10 patients (49). Two cell lines derived from a primary poorly differentiated PeSCC and its corresponding lymph node metastasis were successfully established. To confirm the malignant potential of these two cell lines, the authors injected the tumor cells subcutaneously in SCID (severe combined immunodeficiency) mice. The authors found similar morphological and immunohistological features comparing cell lines with the derived xenograft tumor. Decreased expression of CXCL14 was detected in the cell line derived from the lymph node metastasis, which suggested a mechanism of immune surveillance escape during tumor cell migration to lymph nodes.

Out of 21 penile tumor tissues, one cell line (Pen1) derived from a lymph node metastasis was successfully established and characterized by Chen et al. (50). Pen1 cells were tumorigenic in SCID mice, presented deleterious TP53 mutation, and increased expression of EGFR and PEDN.

We also reported a comprehensive characterization of a cell line and xenograft derived from a verrucous PeSCC, accounting for 2–8% of PeCa cases (51). The cell line and xenografts were comprehensively characterized using immunophenotyping, and large-scale genomic and transcriptomic analyses. Genomic alterations observed in the cell line and xenografts showed high similarity with the parental PeSCC. Interestingly, the tumor generated in the BALB/c nude mice presented a sarcomatoid-like carcinoma phenotype. As a result, this study demonstrated that xenograft PeSCC models must be used with precaution, considering the selection of specific cell populations and anatomical sites where the cells or tumor are implanted.

The first platinum-resistant penile cancer-patient derived xenograft (NOD/SCID/IL2γ-receptor null mice) was recently established and characterized (52). Small animal imaging was used as proxies for therapeutic efficacy providing further output on tumor perfusion and metabolic activity. Humanized mice models are exciting alternatives to testing immune checkpoint blockade (52).

Whole-genomic sequencing assays were performed in four PeSCC cells lines established by Zhou et al. (53). These cell lines were derived from HPV-negative cases and showed tumorigenicity in nude mice. The authors reported variants in ERCC5, TP53, PTH1, CLTCL1, NOTCH2, MAP2K3, CDK11A/B, USP6, ADCH5, BCLAF1, CDKN2A, FANCD2, HRAS, and NOTCH1. Amplifications of MYC
and EGFR and losses of FBXW7, TET2, XPC, and FANCE were also described. A similar portrait of the genomic alterations was observed comparing tumor and the derived cell line. Also, the pathways altered in these cell lines were previously described in PeSCC, including MAPK, Jak-STAT, TGF-beta, Notch, and apoptosis signaling pathway.

Fenner et al. established four cell lines derived from primary PeSCC and lymph node metastasis (54). Invasion and capillary tube formation assays, chemoresponsiveness, and mRNA and protein expression analyses were investigated in these cell lines. The authors described deregulation of RB/E2F1 axis in metastatic cells and concluded that E2F1 is a driver of invasion, lymphatic dissemination, and promotes chemoresistance.

A pivotal study described the generation and characterization of the first genetically engineered mouse models of PeSCC (SA: PB-Cre4 Smad4L/L, ApcL/L, and SAP: PB-Cre4 Smad4L/L, ApcL/L, PtenL/L, mice) (55). The authors showed that a single knock-out model was insufficient to drive penile tumorigenesis, only achieving success by applying Smad4 and Apc co-deletion in the androgen-responsive epithelium of the penis (55). The murine PeSCC presented gene signatures comparable with those described in humans. The single-cell analysis revealed an intratumoral immunosuppressive myeloid cell infiltration in the SA mice. A randomized pre-clinical trial using these models and immune-checkpoint inhibitors with or without targeted therapy showed that tumor eradication was achieved only upon combining different drugs. This study presented a valuable platform for testing and discovering treatment strategies, and results obtained by the authors suggested that combined target therapy and immunotherapy could be used in the treatment of PeSCC patients.

Contemporary protocols

Evolution of the current systemic strategies

Given the low incidence of PeSCC and inadequate centralization of care in the majority of countries, coupled with limited up-to-date expertise of physicians in the disease management, current concepts of systemic treatment are based on the findings of small and mostly retrospective case-series assessing plenty of different regimens (56,57). Thus, Protzel and co-authors reported 18 different chemotherapy regimens used in 91 Germany centers, whereas chemotherapy for PeSCC was performed on average 2.3 times annually per department (58). This ultimately translates into low compliance of treatment decisions with guideline recommendations. Thus, Distler and collaborators demonstrated adherence to the PeSCC guidelines of the European Association of Urology (EAU) for a neoadjuvant, adjuvant and palliative chemotherapy indication in 21%, 26% and 48%, respectively (59). Furthermore, the level of evidence generated from the findings of these trials is attenuated by long periods of assessment with varying class and dose of agents, mode of application, number of courses, characteristics as well as compliance issues of participating patients.

Platinum-based agents, in particular cisplatin, are undoubtedly the mainstay of the current systemic treatment strategies of PeSCC alongside a wide range of other solid neoplasms. One of the most pertinent mechanisms of its anticancer activity is the induction of DNA lesions promoting activation of the DNA damage response and induction of mitochondrial apoptosis and subsequent cell death (60). Hereby, cisplatin resistance can rest upon alterations (I) in processes that predate its binding to DNA and cytoplasmic structures (pre-target resistance), (II) directly related to the molecular damage caused by cisplatin (on-target resistance), (III) in the lethal signaling pathways triggered by such molecular lesions (post-target resistance) and 4) influencing molecular mechanisms not associated with cisplatin-elicited signals (off-target resistance) (61). Notably, cancers with TP53 mutated respond worse to cisplatin than those harboring TP53 wild type as the post-target mode of cisplatin resistance (60). In turn, overactivation of ERBB2, consequently stimulating PI3K/AKT signaling, has been identified as the off-target mechanism of chemoresistance in several malignancies, e.g., non-small cell lung and gastric cancer (62,63). Additionally, PTEN deficiency has been reported to confer cisplatin resistance in PeSCC (55). As outlined above, these molecular alterations are frequently encountered in PeSCC, at least partially elucidating its poor responsiveness to cisplatin and emphasizing the need for effective combination protocols targeting different drug resistance pathways.

The limited antitumor activity of single agents such as cisplatin, bleomycin, and methotrexate was first reported in 1970 (56). However, response rates were minimal, observed at 0–27%, and significant side effects such as bleomycin-associated pulmonary toxicity were observed (56,64-68). These unsatisfactory outcomes paved the way for combination regimens. The increasing body of evidence for efficacy of cisplatin as a combination partner in protocols for SCC of different origins contributed to the systemic
management of PeSCC (56). First informative case-series on the dual systemic therapy of advanced PeSCC originate from the late 1980s and early 1990s. Hussein and collaborators reported on treating six patients (including one metastatic) with recurrent or unresectable disease affecting penis or urethra with a combination of 100 mg/m$^2$ cisplatin on d1 and 5-fluorouracil (5-FU) at a dose of 960 mg/m$^2$/d on d2–6 every 3 to 4 weeks (69). Alopecia was universal, while other toxicities like mucositis, nausea, and vomiting were mild. All patients experienced partial or complete remission and some of them were submitted to surgery and radiotherapy, while survival ranged between 6 and 32 months in this mixed cohort of neoadjuvant and palliative settings. Shammas et al. utilized the same protocol (except for 5-FU dose of 1,000 mg/m$^2$/d) for eight patients in the neoadjuvant setting, observing partial remission in 25% of cases (70). Severe toxicities, including deterioration of renal function in three and sepsis in two patients, hampered common application of this protocol, especially in elderly individuals. Subsequent reports analyzed other dual combinations of cisplatin like those with methotrexate, adriamycin or irinotecan, yielding response rates ranging from 0 to 31% (71,72). In addition, Power and colleagues reported two patients treated with cisplatin 80 mg/m$^2$ on d1 + gemcitabine 1.250 mg/m$^2$ on d1 and d8, both experiencing partial remission (73).

Several triple regimens have been investigated aiming to improve the outcome of PeSCC patients. The first triple-drug protocol was described by Pizzocaro et al. They subjected twelve patients with resected lymph node metastases as well as five patients with fixed inguinal nodes to 12 weekly courses of 15 mg/m$^2$ bleomycin, 1 mg/m$^2$ vincristine, and 30 mg/m$^2$ methotrexate (BVM) in the adjuvant and neoadjuvant indication, respectively (74). With the median follow-up of 42 months, only one out of twelve patients managed in the adjuvant setting relapsed, whereas in two out of five patients treated in the neoadjuvant setting, surgical tumor resection was incomplete during lymphadenectomy leading to death within twelve months. The most critical toxicities included 1x myelosuppression and 2x suspension of the protocol after eight courses due to lung fibrosis. Dexeus and co-authors applied bleomycin 10 mg/m$^2$ on d2–6, methotrexate 200 mg/m$^2$ on d1 and d15, and cisplatin 20 mg/m$^2$ on d2–6 (BMP) in fourteen patients observing a response rate of 72% with a response duration of 6 months (56,75). These treated patients showed mild side effects like hypercalcemia and infection (56,75). The hope arose that this protocol will get the standard of care, but further clinical evidence proved it wrong. Thus, Haas and collaborators reported a 32.5% response rate at the cost of five treatment-related deaths and six cases with life threatening toxic episodes in a prospective, nonrandomized study of forty patients treated with a slightly dose-modified “Dexeus regimen” (76). Similar findings were reported by others using this therapeutic scheme (77,78).

Further efforts were aimed at assessing the combination of a platinum and taxane with a further compound. Pizzocaro et al. subjected six patients to neoadjuvant paclitaxel 120 mg/m$^2$ d1, cisplatin 50 mg/m$^2$ d1–2 and 5-FU 1,000 mg/m$^2$d2–5 (TPF) leading to a response rate of 83.3% and solely grade 2 side effects (79). The landmark phase II study by Pagliaro and co-authors from the MD Anderson Cancer Center in Houston, USA, treated 30 PeSCC patients with four courses of paclitaxel 175 mg/m$^2$ d1, ifosfamide 1,200 mg/m$^2$d1–3, and cisplatin 25 mg/m$^2$d1–3 (TIP) in a prospective, nonrandomized, phase II trial (80). The response rate was 50%, including three men with pathologically complete remission, while the median time to progression was 8.1 months and overall survival of 17.1 months. The regimen was well tolerated without treatment-related deaths, while infection grade 3 was the most common toxicity. In a former report from the same center on a retrospective assessment of 10 men treated in the neoadjuvant setting with different protocols, the only three ones with pN0 on surgical procedure received TIP (81). Selected studies forming a basis for contemporary systemic management of advanced PeSCC are depicted in Table 1.

Neoadjuvant indication

The primary goal of the neoadjuvant treatment is the shrinking of initially as not completely operable classified tumor (inductive therapy) followed by an in sano surgical resection (main therapy). EAU guidelines advocate to offer to PeSCC patients with non-resectable or recurrent lymph-node metastases four cycles of cisplatin- and taxane-based regimen followed by radical surgery (82). This refers mostly to clinically fixed inguinal nodal masses (cN3). Moreover, neoadjuvant chemotherapy may be an option for locally advanced (T4) and ulcerated cancers (82). There is no clear consensus on what regimen should be the standard of care. TIP may be utilized based on the aforementioned data by Pagliaro et al. (80) and Bermejo et al. (81). Recently, Xu and collaborators observed a response rate of 63.2% with docetaxel 75 mg/m$^2$ d1, cisplatin 25 mg/m$^2$d1–3, and ifosfamide 1,200 mg/m$^2$
d1–3 in 19 patients with only one male discontinuing chemotherapy due to severe myelosuppression (83). On the other hand, TPF also provided promising results in a small study by Pizzocaro and co-authors (79). Similarly, using up to three courses of docetaxel 75 mg/m² d1, cisplatin 60 mg/m² d1, and 5-FU 750 mg/m² on d1–5 in 29 men with locally advanced or metastatic PeSCC, Nicholson and colleagues demonstrated a response rate of 36.8% in 19 patients treated with neoadjuvant therapy. However, 65.5% of all men experienced at least one grade 3/4 adverse event (84). A crucial aspect of the neoadjuvant approach is the re-evaluation of response after two cycles and treatment continuation only in responders. A recent meta-analysis by Azizi et al. included 182 patients from 10 studies, in which 66 (36.3%) and 116 (63.7%) males were treated respectively with nontaxane-platinum and taxane-platinum regimens (85). The analysis demonstrated that about 50% of patients with bulky regional lymph node metastases responded to platinum based neoadjuvant chemotherapy as well as 16% of men achieved pathologically complete response, thus substantiating the value of neoadjuvant systemic chemotherapy in bulky (cN2/N3) penile cancer.

**Adjuvant indication**

The adjuvant strategy is scheduled to eradicate clinically inapparent micrometastases after complete surgical treatment. EAU guidelines recommend offering to patients with pN2/N3 disease (metastasis in three or more unilateral or bilateral inguinal nodes, metastasis in pelvic lymph nodes, and extranodal extension of regional lymph node metastasis) after radical lymphadenectomy adjuvant chemotherapy with 3–4 courses of cisplatin, a taxane, and 5-FU or ifosfamide (82). Similar to the neoadjuvant indication, no consensus on the standard of care protocol exists in this setting. Giannatempo et al. treated with adjuvant therapy 19 patients with 3–4 cycles TPF (paclitaxel 120 mg/m² d1 or docetaxel 75 mg/m² d1, cisplatin 75–100 mg/m² d1 + 5-FU 750–1,000 mg/m² d1) achieving remission in 52.6% after a median follow-up of 42 months (86). The 2-year disease-free survival was 36.8% (87). Neutropenia grade 3–4 was the most common toxicity, observed at 21%. O’Reilly and collaborators reported on treating 3 patients with four courses of TIP (paclitaxel 175 mg/m² d1, ifosfamide 1,200 mg/m² d1–3, and cisplatin 20 mg/m² d1–3) without signs of recurrence after 6, 28 and 50 months, respectively (88).

**Table 1** Selected studies of the systemic treatment of advanced penile squamous cell carcinoma (PeSCC)

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<th>Indication</th>
<th>Autor</th>
<th>Number treated</th>
<th>Regimen</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjuvant</td>
<td>Pagliaro et al., J Clin Oncol, 2010</td>
<td>30</td>
<td>4x paclitaxel 175 mg/m² d1, ifosfamide 1,200 mg/m² d1–3 + cisplatin 25 mg/m² d1–3 (TIP)</td>
<td>RR 50%; TTP 8.1 mos.; OS 17.1 mos.</td>
</tr>
<tr>
<td>Neoadjuvant</td>
<td>Pizzocaro et al., Eur Urol, 2009</td>
<td>6</td>
<td>4x paclitaxel 120 mg/m² d1, cisplatin 50 mg/m² d1–2 + 5-FU 1,000 mg/m² d2–5 (TPF)</td>
<td>RR 83.3%</td>
</tr>
<tr>
<td>Neoadjuvant</td>
<td>Xu et al., BMC Cancer, 2019</td>
<td>19</td>
<td>4x docetaxel 75 mg/m² d1, cisplatin 25 mg/m² d1–3 + ifosfamide 1,200 mg/m² d1–3 (TIP)</td>
<td>RR 63.2%</td>
</tr>
<tr>
<td>Adjuvant</td>
<td>Nicolai et al., Clin Genitourin Cancer, 2016</td>
<td>19</td>
<td>3-4x paclitaxel 120 mg/m² d1 or docetaxel 75 mg/m² d1, cisplatin 75-100 mg/m² d1 + 5-FU 750-1,000 mg/m² d1 (TPF)</td>
<td>2-year PFS 36.8%</td>
</tr>
<tr>
<td>Adjuvant</td>
<td>O’Reilly et al., J Clin Oncol, 2013</td>
<td>3</td>
<td>4x paclitaxel 175 mg/m² d1, ifosfamide 1,200 mg/m² d1–3 and cisplatin 20 mg/m² d1–3 (TIP)</td>
<td>PF at 6, 28 and 50 mos.</td>
</tr>
<tr>
<td>Palliative</td>
<td>Zhang et al., Oncotarget, 2015</td>
<td>39</td>
<td>4x docetaxel 75 mg/m² d1, cisplatin 70 mg/m² d1 + 5-FU 500 mg/m²/d d1–5</td>
<td>RR 38.5%; PFS 3 mos.; OS 7 mos.</td>
</tr>
</tbody>
</table>

PeSCC, penile squamous cell carcinoma; RR, response rate; OS, overall survival; TTP, time to progression; PF(S), progression-free (survival); mos.: months; d, day(s); 5-FU, 5-fluorouracil; TIP, taxane, ifosfamide + platinum; TPF, taxane, platinum + 5-FU.

**Palliative indication**

Since there is no cure for visceral disseminated PeSCC, systemic therapy is aimed at prolonging survival and alleviating symptoms. EAU guidelines recommend performing chemotherapy in patients with systemic disease, alluding to better efficacy of cisplatin- and taxane-including
regimens (82). Zhang et al. treated 39 patients with a median of four courses of TPF (docetaxel 75 mg/m² d1, cisplatin 70 mg/m² d1, and 5-FU 500 mg/m²/d d1–5) (89). The response rate was as high as 38.5%, progression-free survival was of 3 months, and overall survival of 7 months. The most frequent adverse events of grade 3 or higher were neutropenia (33%). In the abovementioned study of Nicholson and colleagues, the response rate to TPF in 7 patients with distant metastasis was 42.9% (84). In line with neoadjuvant and adjuvant indications, both TPF and TIP appear to be reasonable options as the first-line treatment.

Upon progression, there are virtually no consistent data on systemic treatment options. Benefit has been reported in small case-series and casuistries for several drugs, e.g., panitumumab, cetuximab, docetaxel, paclitaxel, sunitinib, and sorafenib (90-93). Depending on availability, treating patients in clinical trials is currently a preferable option.

**Ongoing clinical trials and future directions**

Several clinical trials are explicitly recruiting patients with advanced PeSCC (94). A phase 2 study is set to investigate the efficacy of the PD-L1 inhibitor avelumab in locally advanced or metastatic PeSCC, in which patients are judged unfit for or have progressed during/after platinum-based chemotherapy (NCT03391479). PULSE phase 2 trial is assessing the effect of avelumab maintenance strategy in patients who are in response or with stable disease after first-line platinum containing polychemotherapy (NCT03774901). AFU-GETUG 25 phase 2 trial is evaluating the value of TIP in the neoadjuvant or adjuvant setting of the multimodal approach by bilateral lymphadenectomy and chemotherapy (NCT02817958). InPACT is a phase 3 study comparing neoadjuvant chemotherapy (TIP) followed by inguinal lymphadenectomy or neoadjuvant chemoradiotherapy (radiotherapy with concurrent cisplatin) followed by inguinal lymphadenectomy to the standard lymphadenectomy (NCT02305654). A number of basket trials are recruiting PeSCC patients as well investigating different protocols, e.g., PD-1 inhibitor pembrolizumab (NCT02721732), c-Met inhibitor cabozantinib + PD-1 blocker nivolumab + CTLA-4 inhibitor ipilimumab (NCT02496208), DNA Plasmid-encoding Interleukin-12/HPV DNA Plasmids Therapeutic Vaccine INO-3112 combined with PD-L1 blocker durvalumab in HPV-associated cancers (NCT03439085).

Along with the advent of the evidence on molecular machinery and causal drivers of PeSCC progression, precision medicine based on personalized genomic, transcriptional, or protein expression profiling might be materialized in this condition in the future. A number of molecular alterations have been recently identified in PeSCC, e.g., related to HRAS, mTOR, VEGF, NOTCH1, and PIK3CA, in addition to a high rate of PD-L1 expressing tumor cells as well as tumor-infiltrating leukocytes (2,95). These insights form a basis for the discovery of actionable targets for drug repurposing or novel targeted agents and transition of their testing through the pipeline from cell cultures over xenograft models into clinical trials.

**Conclusions**

Current systemic protocols for the treatment of advanced PeSCC are characterized by limited efficacy combined with considerable side effects. Fortunately, knowledge of molecular machinery causally involved in its tumorigenesis is expanding. Further joint research efforts between basic and clinical researchers are warranted to realize the concept of individualized medicine in this rare disease.

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