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Highlights in genitourinary cancers

T. Vermassen, PhD, S. Rottey, MD, PhD

Department of Medical Oncology and Drug Research Unit Ghent, Ghent University Hospital, Ghent, Belgium

From June 1st till June 5th, Chicago was host for the 55th annual ASCO meeting. This report will highlight the most important studies concerning genitourinary cancers presented during the meeting.

PROSTATE CANCER (PC)

Several new treatment modalities for PC were presented on ASCO. An overview is given in *Table 1*.

The phase III trial GETUG-AFU 16 explored the addition of androgen deprivation therapy (ADT) to salvage radiotherapy (RT) after biochemical recurrence following prostatectomy. As RT + ADT resulted in an increased metastatic-free survival (MFS) after 9 years of follow up, standard addition of ADT to salvage RT could postpone aggressive treatment without increased toxicity or decline in quality-of-life (QoL).¹

The phase III study ENZAMET determined the possible addition of docetaxel or abiraterone acetate to testosterone suppression in metastatic hormone-sensitive prostate cancer (mHSPC) patients to improve overall survival (OS). Interim survival data demonstrate a significantly improved OS by adding enzalutamide to SOC for mHSPC.² Also the phase III trial TITAN assessed the addition of the androgen receptor (AR) inhibitor apalutamide to ADT in mHSPC. A clear improvement in progression-free survival (PFS) and OS were observed, with manageable toxicity profile and no changes in QoL.³ Both the ENZAMET and TITAN study indicate a clear shift of AR inhibitors for treatment of hormone-sensitive PC. Numerous trials assessed new treatment options for non-metastatic and metastatic castrate-resistant prostate cancer (nmCRPC and mCRPC). The phase III trial ARAMIS evaluated the use of the AR antagonist darolutamide in the nmCRPC setting. Darolutamide clearly prolongs MFS, is well tolerated, maintains QoL, and delays worsening of pain and disease-related symptoms compared to placebo.4

The phase II TAXOMET study reported no clinically meaningful addition of metformin to docetaxel for treatment of mCRPC patients. Data from the STAMPEDE trial (SOC + metformin) is expected.

The Alliance A031021 phase III trial compared the combination of enzalutamide + abiraterone acetate versus enzalutamide only. The combination showed no benefit in OS with more treatment-related AEs. The combination of enzalutamide + abiraterone acetate is therefore not recommended.⁶ The phase Ib/II trial KEYNOTE-365 explored the possibility of administering pembrolizumab + enzalutamide in patients who progressed on abiraterone acetate within six months. Promising results were observed (doubling of objective response rate [ORR] compared to pembrolizumab in monotherapy) indicating the possible role of immune checkpoint inhibition (ICI) in the mCRPC setting. The phase III trial KEYNOTE-641 is currently ongoing.⁷

The phase II TOPARP-B trial assessed the use of the poly(AD-P)-ribose polymerase inhibitor olaparib in mCRPC patients with DNA damage repair alterations. The trial demonstrated antitumor activity, especially in patients with *BRCA1/2* loss, *PALB-2* mutations and *ATM* mutations.⁸

Additionally, another phase II trial evaluated cabazitaxel versus enzalutamide or abiraterone acetate in poor prognosis mCRPC patients. It was found that cabazitaxel gives high clinical benefit for poor risk mCRPC patients, although no gain in OS was observed. ctDNA fraction, AR amplification and *TP53* mutations proved to have prognostic value although larger study groups are needed to confirm this

Please send all correspondence to: S. Rottey, MD, PhD, Department of Medical Oncology, Ghent University Hospital, C. Heymanslaan 10, 9000 Ghent, Belgium, Tel: +32 (0)9/332 26 92, E-mail: Sylvie.Rottey@UGent.be

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follow up; mHSPC, metastatic hormone-sensitive prostate cancer; mMFS, median metastatic-free survival; mo, months; NR, not reached; mOS, median overall survival; mPFS, median progression-free survival; (n)mCRPC, (non-)metastatic castrate-resistant prostate cancer; ORR, objective response rate; Pbo, placebo; Pembro, pembrolizumab; PSADT, PSA doubling time; RP, radical prostatectomy; RT, radiotherapy; TS: testosterone suppression. AA, abiraterone acetate; ADT, androgen deprivation therapy; AE, adverse event; Apa, apalutamide; BCR, biochemical recurrence; Caba, cabazitaxel; Enza, enzalutamide; HR, hazard ratio; mFU, median

TABLE 1. New treatment modalities for prostate cancer.TrialGETUG-AFU16ENZAMETTIReference121PhaseIIIIIIIII	GETUG-AFU16 1 III	enzamet 2 III	cancer. TITAN 3	ARAMIS	= 5	A031021		KEYNOTE-365 7 Ib/II	KEYNOTE-365 TOPARP-B 7 8 Ib/II II mCRPC with
Type of patients	BCR after RP	mHSPC	mHSPC	nmCHPC (PSADT ≤ 10 mo)	MCRPC		MCRPC	mCRPC (post AA)	
Number of patients	743	1125	1052	1509	66		1311	1311 69	69
Randomisation	1:1	1:1	1:1	2:1	1:1		1:1	1:1 –	
Therapy	RT + ADT vs. RT	TS + Enza vs. TS + non-steroidal ADT	ADT + Apa vs. ADT + Pbo	ADT + daroluta- mide vs. ADT + Pbo	DOCE + metfor- min vs. Doce + Pbo	etfor- Pbo	enza + AA Pbo vs. Enza		Enza + AA vs. Enza
mFU	112 mo	34.0 mo	22.6 mo		41.1 mo		1		I
mMFS	NR vs. NR HR = 0.73 (0.54 - 0.98)	I	I	40.4 vs. 18.4 mo HR = 0.41 (0.34 - 0.50)	I		1	1	
mPFS	NR vs. 108 mo HR = 0.54 (0.43 - 0.68)	NR vs. 27 mo HR = 0.40 (0.33 – 0.49)	NR vs. 22.1 mo HR = 0.48 (0.39 – 0.60)	1	7.4 vs. 5.6 mo	no	52.2 vs. 20.7 mo HR = 0.85 (0.74 - 0.97)		52.2 vs. 20.7 mo HR = 0.85 (0.74 – 0.97)
mOS	NR vs. NR HR = 0.93 (0.63 - 1.39)	NR vs. NR HR = 0.67 (0.52 - 0.86)	NR vs. NR HR = 0.67 (0.51 - 0.89)	NR vs. NR HR = 0.71 (0.50 - 0.99)	24.6 vs. 19.7 mo	no	mo 34.2 vs. 32.5 mo HR = 0.90 (0.78 - 1.05)		34.2 vs. 32.5 mo HR = 0.90 (0.78 - 1.05)
Clinical benefit rate	I	I	I		I		I	1	
PSA decline > 50%	1	I	I		66 vs. 63 %		80 vs. 82 %	80 vs. 82 % 26 %	vs. 82 %
ORR	I	I	I		28 vs. 28 %		I	- 20 %	
Grade 3/4 AEs	No increase in toxicity	I	42 vs. 41 %		I		69 vs. 56 %	69 vs. 56 % 41 %	vs. 56 %

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finding.9

Finally, the phase III trial CABADOC determined the patient preference between cabazitaxel and docetaxel for firstline chemotherapy in mCRPC. Although cabazitaxel and docetaxel have similar efficacy when used as first-line treatment option, more patients prefer cabazitaxel. Preferable choice was mostly influenced by fatigue, patient-defined QoL, hair loss, and pain.¹⁰

RENAL CELL CARCINOMA (RCC)

It remains a constant point of discussion when to start systemic therapy in metastatic RCC (mRCC) patients, especially in patients with low tumor burden or slow growing disease. The Canadian Kidney Cancer information system identified 1711 patients who immediately started systemic therapy (N=848); started systemic therapy ≥ 6 months after diagnosis of mRCC (N=370) or never received systemic therapy (N=493). Five year-OS was significantly lower for patients who immediately started systemic therapy (32.1 versus 70.2%). After adjusting for IMDC risk criteria and age, both OS (HR 0.46, 0.38-0.56) and time to treatment failure (HR 0.79, 0.69-0.92) were greater for delayed versus immediate systemic treatment. These data suggest that a subset of patients may be safely observed without immediate initiation of systemic therapy, which could be explained by the fewer metastatic sites and increased performance of metastasectomies in this patient group. Prospective validation in the contemporary immunotherapy era is required.¹¹

Next, several treatment modalities for mRCC were presented at ASCO. An overview is given in *Table 2*.

The phase III trial E2810 evaluated the effect of pazopanib on MFS in mRCC treatment-naïve patients with no evidence of disease following metastasectomy. The primary end point was not reached and adjuvant pazopanib in this patient cohort is thus not recommended.¹²

The phase III CARMENA trial previously indicated that cytoreductive nephrectomy (CN) is not advised in mRCC. Updated results strengthen this statement. However, it was shown that patients with only 1 IMDC risk criteria could still benefit from CN.¹³

A phase II trial by Gao *et al.* evaluated the benefit of concomitant CN or metastasectomy in mRCC patients receiving firstline ICI. The authors suggest that ICI plus concomitant CN or metastasectomy is safe and shows promising clinical utility. Furthermore, response to therapy and survival outcome might be correlated to several biomarkers, such as CD8 tumor infiltrating lymphocytes and tumor IFN.¹⁴

The phase II CheckMate 920 study determined the clinical efficacy of ICI in patients with brain metastases. The current results show encouraging efficacy results with safety profile

comparable to previous reported studies.¹⁵

Finally, several subanalyses of large phase III studies were presented in which the effect of ICI on sarcomatoid mRCC and IMDC intermediate and poor risk mRCC were assessed. IMmotion 151, CheckMate 214 and KEYNOTE-426 all showed high benefit from ICI for patients with sarcomatoid features and intermediate and poor risk patients.¹⁶⁻¹⁸

The fact that sarcomatoid mRCC respond well to ICI can be partly explained by the retrospective analysis done by Bakouny *et al.* After performing next-generation sequencing on sarcomatoid and rhabdoid mRCC tumors, analysis showed that genomic alterations in *BAP1* were significantly more frequent in sarcomatoid and rhabdoid mRCC (25 vs. 4.3%) while other genomic alterations and tumor mutational burden were similar. This could account for the fact that sarcomatoid and rhabdoid mRCC tumors have better outcomes on ICIs compared to non-ICI-based therapies.¹⁹

In addition, patient reported outcomes from the IMmotion 150 suggested that atezolizumab, alone or with bevacizumab, maintained daily function with minimal symptom interference versus sunitinib.²⁰

UROTHELIAL CARCINOMA (UC)

Numerous novel therapies for treatment of (metastatic) urothelial carcinoma (mUC) were presented at ASCO. An overview is given in *Table 3*.

First, the most ideal adjuvant therapy following cystectomy in patients with locally advanced disease was determined. Comparison between adjuvant RT or chemotherapy proved comparable MFS, although local control is improved in the RT arm. Based on this study, this treatment option could be offered for patients unfit or unwilling to receive chemotherapy.²¹

The CALGB 90601 phase III study assessed the added value of bevacizumab to chemotherapy in treatment-naïve mUC. No OS benefit was shown. A small gain in PFS was observed, although not clinically significant. Bevacizumab has therefore no place in first-line therapy.²²

The HCRN GU14-180 phase II trial explored the role of maintenance ICI in patients who are stable after first-line chemotherapy. Maintenance ICI proved effective and prolonged PFS. Further validation is even though still required to verify if maintenance ICI "deepens" responses achieved with first-line chemotherapy.²³

Response to ICI may be dampened by *FGFR3* mutations. The phase Ib/II FIERCE-22 trial therefore explored the efficacy of the combination of the FGFR3 inhibitor vofatamab and ICI. The combination seems well tolerated and prolongs PFS, especially in patients with wild type *FGFR3*. Further investigation is ongoing.²⁴

Atezo, atezolizumab; axi, axitinib; bev, bevacizumab; HR, hazard ratio; IMDC, International Metastatic RCC Database Consortium; ipi, ipilimumab; mFU, median follow up; mMFS, median metastatic-free survival; mo, months; mOS, median overall survival; mPFS, median progression-free survival; nivo, nivolumab; NR, not reached; ORR, objective response rate; pazo, pazopanib; pbo, placebo; pembro,

Grade 3/4 AEs	ORR	SOW	mPFS	mMFS	mFU	Therapy	Randomisation	Number of patients	Type of patients	Phase	Reference	Trial
I	I	NR vs. NR HR = 2.65 (1.02 - 6.9)	I	17.3 vs. 14.2 mo HR = 0.85 (0.55 - 1.31)	30 mo	Pazo vs. pbo	1:1	129	Treatment-naïve (no evidence of disease following metastasec- tomy)	=	12	E2810
I	I	15.6 vs. 19.8 mo HR = 0.93 (0.76 – 1.15)	I	I	61.5 mo	Sun + CN vs. sun	1:1	450	Treatment-naïve	=	13	CARMENA
All: 38 vs. 42 vs. 47 %	+ surgery: 86 vs. 89 vs. 69 % - surgery: 55 vs. 44 vs. 43 %	All: NR vs. NR vs. NR	+ surgery: 17.3 vs. 7.6 vs. 8.9 mo - surgery: 14.5 vs. 7.6 vs. 7.5 mo	I	24.6 mo	Nivo ± surgery vs. nivo + bev ± sur- gery vs. nivo + ipi ± surgery	2:3:2	105	Treatment-naïve	=	14	NCT02210117
21 %	28.6 %	Υ. Έ	9.0 mo	I	6.5 mo	Nivo + ipi	I	28	Treatment-naive (asymptomatic brain metastases)	=	15	CheckMate 920
40 vs. 49 %	49 vs. 14 %	NR vs. 15.0 mo HR = 0.56 (0.32 - 0.96)	8.3 vs. 5.3 mo HR = 0.52 (0.34 – 0.79)	I	I	Atezo + bev vs. sun	1:1	142/863	Treatment-naïve (sarcomatoid)	=	10	IMmotion 151
46 vs. 40 %	56.7 vs. 19.2 %	31.2 vs. 13.6 mo HR = 0.55 (0.33 - 0.90)	8.4 vs. 4.9 mo HR = 0.61 (0.38 - 0.97)	I	30 mo	Nivo + ipi vs. sun	1:1	112/1096	Treatment-naïve (sarcomatoid and IMDC intermediate / poor risk)	=	17	CheckMate 214
I	IMDC: 55.8 vs. 29.5 % sarcomatoid: 58.8 vs. 31.5 %	IMDC: NR vs. NR HR = 0.52 (0.37 - 0.74) sarcomatoid: NR vs. NR HR = 0.58 (0.21 - 1.59)	IMDC: 12.6 vs. 8.2 mo HR = 0.67 (0.53 - 0.85) sarcomatoid: NR vs. 8.4 mo HR = 0.54 (0.29 - 1.00)	I	I	Pembro + axi vs. sun	1:1	IMDC: 592/861 sarcomatoid: 105/861	Treatment-naïve (sarcomatoid / IMDC intermediate / poor risk)	=	18	KEYNOTE-426

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TABLE 3. New treatment modalities for advanced urothelial carcinoma.	ent modalities for adv	anced urothelial carc	sinoma.				
Trial	NCT01734798	CALGB 90601	HCRN GU14-182	FIERCE-22	NCT03333616	NCT03507166	EV-201
Reference	21	22	23	24	25	26	27
Phase	=	=	=	II/qI	=	=	=
Type of patients	Chemo-naïve, local disease after cystec- tomy	Treatment-naïve (> 12 mo since adjuvant chemotherapy)	First line chemo-pre- treated patients with stable disease	≥ 1 prior chemo- therapy or < 12 mo since adjuvant chemo- therapy	Variant histologies, treatment-naïve or pretreated (no ICI)	HER2+, pretreated (≥ 1 prior systemic therapy)	Pretreated (prior plati- num chemotherapy and ICI)
Number of patients	123	506	107	7/28	19	43	128
Randomisation	2:1			I	I	I	I
Therapy	RT vs. adjuvant chemo- therapy	Chemotherapy + bev vs. chemotherapy + pbo	Pembro vs. pbo	Vofatamab + pembro	Nivo + ipi	RC48-ADC	Enfortumab vedotin
mFU	I	46.2 mo	14.7 mo	I	3.6 mo		4.6 mo
mMFS	HR = 0.65 (0.35 - 1.19)	I	I	I	I	I	I
mPFS	I	7.7 vs. 6.6 mo HR = 0.79 (0.66 - 0.95)	5.4 vs. 3.2 mo HR = 0.64 (0.41 - 0.98)	К	3.8 mo	6.9 mo	5.8 mo
SOm	HR = 0.94 (0.52 – 1.69)	14.5 vs. 14.3 mo HR = 0.87 (0.72 - 1.06)	I	I	КN	Ш	11.7 mo
ORR	I	40.4 vs. 33.0 %	22 vs. 12 %	36 %	37 %	51.2 %	44 %
Grade 3/4 AEs	8 vs. 2 % (late GI toxicity)	83.5 vs. 80.7 %	53 vs. 35 %	1	16 %	I	I

Bev, bevacizumab; HR, hazard ratio; ICI, immune checkpoint inhibition: ipi, ipilimumab; mFU, median follow up; mMFS, median metastatic-free survival; mOS, median overall survival; mPFS, median progression-free survival; nivo, nivolumab; NR, not reached; ORR, objective response rate; pbo, placebo; pembro, pembrolizumab; RT, radiotherapy.

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KEY MESSAGES FOR CLINICAL PRACTICE

- 1. A clear shift is seen towards modern AR inhibitors for treatment of hormone-sensitive PC.
- 2. Combinations of docetaxel plus metformin or enzalutamide plus abiraterone are not advised for treatment of mCRPC.
- 3. Pembrolizumab plus enzalutamide show promising results in mCRPC (although phase III data are awaited).
- 4. Genetic profiling will play a role in determining the most optimal treatment option for mCRPC.
- 5. Active surveillance remains a valid option for mRCC, especially in patients with low tumor burden.
- 6. Cytoreductive nephrectomy is only recommended in patients with low risk (IMDC 1) and might be plausible in patients receiving first-line ICI.
- 7. ICI proves effective for mRCC patients diagnosed with asymptomatic brain metastases.
- **8.** Sarcomatoid mRCC patients respond well to ICI, probably due to the genomic alterations (especially *BAP1* mutations) that are associated with this histologic feature.
- **9.** Adjuvant RT for locally advanced mUC is a possible option for patients unfit or unwilling to receive chemotherapy.
- 10. Bevacizumab is not to be given in addition to first-line chemotherapy in mUC.
- **11.** Maintenance ICI might "deepen" the responses achieved with first-line chemotherapy in mUC results of first line combination trials are awaited.
- **12.** Combination of ICI and FGFR3 inhibitors might increase the ORR in mUC due to the inhibition of the dampening effect created by *FGFR3* mutations.
- 13. mUC of variant histologies can respond to ICI and show a desirable safety profile.
- **14.** Novel treatment options are coming which show efficacy in mUC patients who progressed on first-line chemotherapy and second-line ICI (enfortumab vedotin and FGFR inhibition are the most important approaches in this setting).

Patients with mUC of variant histologies have poor outcomes. A phase II trial was conducted to assess the use of ICI in this patient group. ICI showed clear efficacy with desirable safety profile. Further exploration of ICI in this patient population is therefore warranted.²⁵

Despite the use of ICI in mUC, the question remains which treatment to choose after progression on ICI. Two phase II trials were reported exploring this statement. RC48-ADC, an anti-HER2 antibody-drug conjugate, proved clinically meaningful in HER2+ patients pretreated with ICI (and chemotherapy).²⁶ Next, enfortumab vedotin, an antibody-drug conjugate targeting Nectin-4, proved effective for patients who progressed after chemotherapy and ICI.²⁷

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