

# The First Slovak Experience with Second-line Vinflunine in Advanced Urothelial Carcinomas

Prvá slovenská skúsenosť s použitím vinflunínu v druhej línii liečby pokročilých urotelových karcinómov

Palacka P., Mego M., Obertova J., Chovanec M., Sycova-Mila Z., Mardiak J.

2<sup>nd</sup> Oncology Department, Medical School of Comenius University in Bratislava and National Cancer Institute, Bratislava, Slovak Republic

## Summary

**Background:** Based on the results of phase III trial, vinflunine was approved by European Medicines Agency in 2010 as second line treatment of advanced urothelial cancer in patients with good performance status (ECOG 0–1). The objective of this prospective observational study was to assess vinflunine treatment of advanced urothelial cancer patients in terms of progression free survival and overall survival, and to evaluate vinflunine toxicity. **Patients and Methods:** From April 2011 to June 2014 a total of 16 patients (100%) with advanced urothelial cancer were treated with vinflunine. The median age was 62 years (range 43–80) and the median Karnofsky index was 90% (range 80–100%). Thirteen patients (81.25%) had urothelial bladder cancers, two patients (12.50%) suffered from urothelial cancers of ureter, and one patient (6.25%) had urothelial cancer of unknown origin (histology was obtained from liver metastasis). Histologically, all the lesions were grade 3 tumors (100%). The number of metastatic sites ranged from 1–4 (median 3). **Results:** The effect of treatment was evaluated in accord with RECIST: two patients (12.50%) obtained partial remission, three (18.75%) stabilization, eight patients (50.00%) progressed, and treatment was suspended in one case at patient's request. Vinflunine toxicity grade 3–4 included neutropenia in six patients (37.50%), leukopenia in four patients (25.00%), anemia in one patient (6.25%), constipation in three patients (18.75%), and febrile neutropenia in one patient (6.25%). Median overall survival was 5.2 months (95% CI 3.4–8.8) and median progression-free survival was 2.3 months (95% CI 2.1–3.2). **Conclusion:** This study summarizes the first Slovak experience with vinflunine therapy. Our data confirmed the efficacy of vinflunine and its acceptable toxicity in the treatment of patients with advanced urothelial cancer previously treated with a platinum-based regimen.

## Key words

advanced urothelial cancer – vinflunine – progression-free survival – overall survival – side effects

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MUDr. Patrik Palacka, PhD.  
MPH, MBA

2<sup>nd</sup> Oncology Department  
Medical School of Comenius  
University in Bratislava and National  
Cancer Institute  
Klenova 1  
833 10 Bratislava 3  
Slovak Republic  
e-mail: info@onkol.sk

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## Súhrn

**Úvod:** EMA (European Medicines Agency) schválila vinflunín v roku 2010 do 2. línie liečby pacientov s pokročilými urotelovými karcinómami v dobrom výkonnostnom stave (ECOG 0–1). Cieľom tejto prospektívnej observačnej štúdie bolo vyhodnotiť liečbu pacientov s pokročilými urotelovými karcinómami vinflunínom z pohľadu prežívania bez progresie, celkového prežívania a toxicity. **Pacienti a metódy:** V období apríl 2011–jún 2014 sme liečili vinflunínom 16 pacientov (100 %) s pokročilými urotelovými karcinómami. Medián veku bol 62 rokov (rozsah 43–80), medián Karnofského indexu bol 90 % (rozsah 80–100 %). Trinásť pacientov (81,25 %) malo urotelové karcinómy močového mechúra, dvaja pacienti (12,50 %) urotelové karcinómy močovodu a jeden pacient (6,25 %) urotelový karcinóm neznámeho pôvodu (histológia bola získaná z pečenej metastázy). Všetky primárne tumory boli stupňa 3 (100 %). Počet metastatických miest bol v rozmedzí 1–4 (medián počtu 3). **Výsledky:** Účinok liečby bol hodnotený v súlade s kritériami RECIST: dvaja pacienti (12,50 %) dosiahli parciálnu remisiu, traja pacienti (18,75 %) stabilizáciu, osem pacientov (50 %) progredovali priamo na liečbe a v jednom prípade bola terapia predčasne ukončená na žiadosť pacienta. Toxicita vinflunínu stupňa 3–4 zahrňovala neutropéniu u šiest pacientov (37,50 %), leukopéniu u štyroch chorých (25,00%), anémiu u jedného pacienta (6,25 %), zápchu u troch chorých (18,75 %) a febrilnú neutropéniu u jedného pacienta (6,25 %). Medián celkového prežívania bol 5,2 mesiacov (95% CI 3,4–8,8), medián prežívania bez progresie 2,3 mesiace (95% CI 2,1–3,2). **Záver:** Táto štúdia predstavuje prvú slovenskú skúsenosť s liečbou vinflunínom. Naše údaje potvrdzujú účinnosť liečby pacientov s pokročilými urotelovými karcinómami predliečenými platínoým režimom vinflunínom s akceptovateľnou toleranciou.

## Kľúčové slová

pokročilý urotelový karcinóm – vinflunín – prežívanie bez progresie – celkové prežívanie – nežiaduce účinky

## Introduction

Bladder cancer is the most common malignancy of the urinary system. In 2007, bladder cancer represented 3.9% of all malignant tumours (n = 511) in Slovak Republic (in comparison to 1.827 cases (5%) diagnosed in Czech Republic at that time), the standardized incidence was 15.0/100,000 (22.4/100,000 in Czech Republic), and mortality in men progressed to 5.37/100,000 (191 cases) (5.9/100,000 (507 cases) in Czech Republic). In both countries, slow, yet steady increase of incidence and mortality was registered in the last three decades. Just recently, mortality began to display a tendency towards stabilization [1].

MVAC regimen (including methotrexate, vinblastine, doxorubicin, cisplatin) is considered to be the standard of first-line treatment in patients with advanced urothelial bladder carcinoma [2,3]. Dose-dense MVAC with granulocyte colony-stimulating factor (G-CSF) support increased complete response rate and improved progression-free survival (PFS) in phase III studies, however, no overall survival (OS) improvement was achieved [4,5]. Another phase III study [6] showed GC (gemcitabine and cisplatin) non-inferiority to MVAC with no significant difference in response rates (49 vs. 46%, respectively), time to progression (7.4 vs. 7.4 months,

respectively), or OS (13.8 vs. 14.8 months, respectively). Grade 3–4 side effects were less frequent in patients treated with GC. Five-year survival rate was comparable in both groups of patients (15 vs. 13 %, respectively) [7]. Due to its comparable efficacy and better side-effect profile, GC became a standard regimen used in the treatment of advanced bladder cancer in majority of EU countries.

There was not any standard approach for second-line treatment of advanced bladder cancer established until recently. Many cytostatics were tested in single-agent setting (phase II studies, Tab. 1) e.g. gemcitabine [8,9], paclitaxel [10], ifosfamide [11,12],

Tab. 1. Monochemotherapy in the second-line treatment of advanced bladder cancer.

Study	Regime	Phase	n	RR (%)	TTP (months)	OS (months)
Lorusso et al [8]	gemcitabine	II	35	23	3.8	5.0
Albers et al [9]	gemcitabine	II	30	11	4.9	8.7
Vaughn et al [10]	paclitaxel	II	31	10	2.2	7.2
Pronzato et al [11]	ifosfamide	II	20	5	nr	nr
Witte et al [12]	ifosfamide	II	56	20	2.5	5.5
McCaffrey et al [13]	docetaxel	II	20	13	nr	9.0
Sweeney et al [14]	pemetrexed	II	47	28	2.9	9.6
Dreicer et al [15]	ixabepilone	II	45	12	2.7	8.0
Bellmunt et al [25]	vinflunine	III	370	9	3.0	6.9

n – number of patients, RR – response rate, TTP – time to progression, OS – overall survival, nr – not reached

**Tab. 2. Combination chemotherapy in the second-line treatment of advanced bladder cancer.**

Study	Regime	Phase	n	RR (%)	OS (months)
Krege et al [16]	docetaxel + ifosfamide	II	22	25	4.0
Lin et al [17]	gemcitabine + ifosfamide	II	23	22	4.8
Bellmunt et al [18]	methotrexate + paclitaxel	II	20	32	5.0
Sternberg et al [19]	gemcitabine + paclitaxel	II	41	60	14.4
Fechner et al [20]	gemcitabine + paclitaxel	II	27	44	13.0
Vaishampayan et al [21]	paclitaxel + carboplatin	II	44	16	6.0
Pagliari et al [22]	ifosfamide + gemcitabine + cisplatin	II	49	41	nr
Chen et al [23]	gemcitabine + docetaxel + carboplatin	I/II	20	45	nr
Tu et al [24]	paclitaxel + cisplatin + methotrexate	II	25	40	nr

n – number of patients, RR – response rate, OS – overall survival, nr – not reached

docetaxel [13], pemetrexed [14], and ixabepilone [15] with response rates (RR) ranging from 5 to 28% (ifosfamid and pemetrexed, respectively), time to progression (TTP) 2.2–4.9 months (paclitaxel and gemcitabine, respectively), and OS from 5.0 months to 9.6 months (gemcitabine and pemetrexed, respectively). Nevertheless, the improvement of both RR and OS was reached mainly with combination chemotherapy (Tab. 2) [16–24]. Particularly, combination of gemcitabine and paclitaxel [19] showed 60% RR and OS of 14.4 months.

Vinflunine (VFL) is a third-generation microtubule inhibitor of the vinca alkaloid class with low and reversible affinity to tubuline. Prior to its clinical evaluation, there was a high anti-tumour activity proven *in vitro*. In a phase III study [25], a total of 370 patients were randomly assigned either to VFL plus best supportive care (BSC),  $n = 253$  or BSC alone,  $n = 117$ . In the response-evaluable population ( $n = 357$ ), the median OS was significantly higher in patients with VFL + BSC in comparison to the BSC group (6.9 vs. 4.3 months, respectively,  $p = 0.040$ ). Overall response rate, disease control, and PFS were all favouring VFL + BSC on a statistically significant level ( $p = 0.006$ ,  $p = 0.002$  and  $p = 0.001$ , respectively). Moreover, the advantage of vinflunine treatment is a predictable and manageable noncumulative toxicity (25%), comprising mainly neutropenia (50%), anemia (19%), fatigue (19%),

constipation (16%) and febrile neutropenia (6%).

The objective of this prospective observational study was to assess the role of vinflunine in the treatment of advanced urothelial cancer in terms of PFS and OS. Another goal was to evaluate toxicity profile of vinflunine, and to compare the results with available literature. This study introduces the first experience with vinflunine therapy in Slovak Republic.

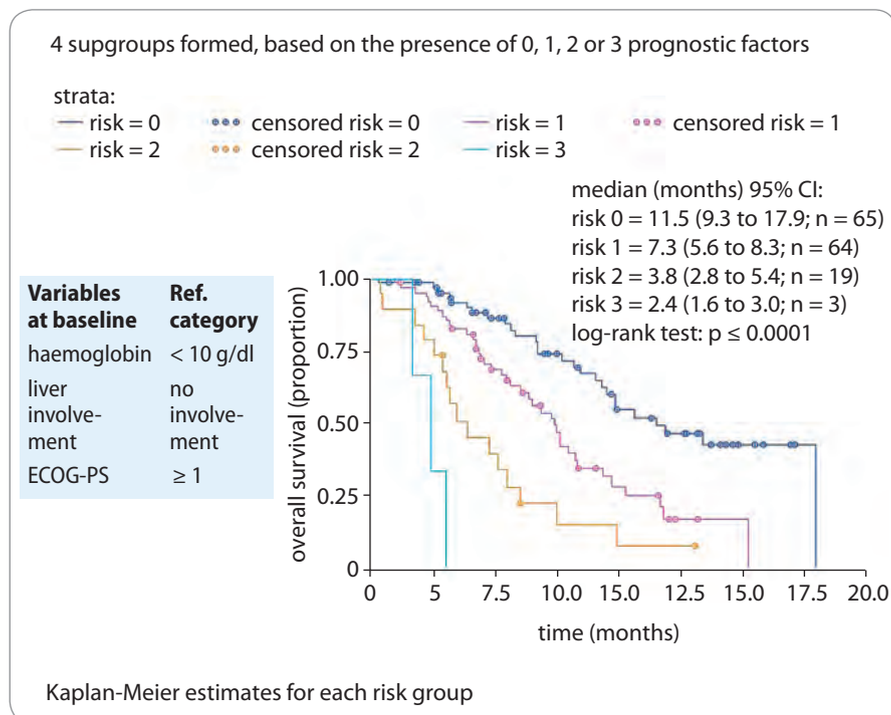
### Characteristics of patients

From April 2011 to June 2014, 16 patients (100%) with advanced urothelial cancer (women:  $n = 3$ , 18.75%; men:  $n = 13$ , 81.25%) were administered vinflunine as second-line therapy at the 2<sup>nd</sup> Oncology Department, Medical School of Comenius University in Bratislava and National Cancer Institute. All patients underwent prior treatment with GC (gemcitabine and cisplatin). The median age was 62 years (range 43–80), and the median Karnofsky performance status at the beginning of treatment was 90% (range 80–100%). Thirteen patients (81.25%) had urothelial bladder cancers, two patients (12.50%) suffered from urothelial cancers of ureter, and one patient (6.25%) had urothelial cancer of unknown origin (histology was obtained from liver metastasis). Histologically, all primary lesions were grade 3 tumors (100%). The number of metastatic sites ranged from 1–4 (median 3). There were several poor prognostic factors

identified in the second-line (Graph 1) treatment settings, such as hemoglobin  $\geq 10$  g/dl in four patients (25.0%), liver involvement in seven patients (43.8%), and ECOG-PS  $\geq 1$  in four patients (25.0%).

### Results

All the patients were administered vinflunine monotherapy in dose of 320 mg/m<sup>2</sup> every three weeks, the number of cycles ranged from 2 to 14 (median 4), and total vinflunine dose was  $2,473.33 \pm 2,108.48$  mg. The tumor response was evaluated after four, eight and twelve cycles in accord with the RECIST [26]: two patients (13.33%) achieved partial response, three patients (20.00%) obtained stabilization, eight patients (53.33%) progressed on the treatment, and therapy was terminated in one case at patient's request, hence was not included in the analysis. The objective response rate was 13.33%. Grade 3 or 4 toxicities associated with vinflunine administration included neutropenia in six patients (37.50%), leukopenia in four patients (25.00%), one patient experienced anemia (6.25%), constipation occurred in three patients (18.75%), and febrile neutropenia (FN) in one patient (6.25%) with a need of further granulocyte colony-stimulating factor (G-CSF) pegfilgrastim prevention every 3 weeks, 48 hours after vinflunine application. Subsequently, no FN events were observed. In the study population, the median PFS (Graph 2) was



Graph 1. Prognostic factors in second-line treatment of advanced bladder cancer [27].

2.3 months (95% CI 2.1–3.2), and the median OS (Graph 3) was 5.2 months (95% CI 3.4–8.8). At the time of final analysis only one patient remained alive; the median follow-up was 5.2 months (95% CI 0.6–16.3).

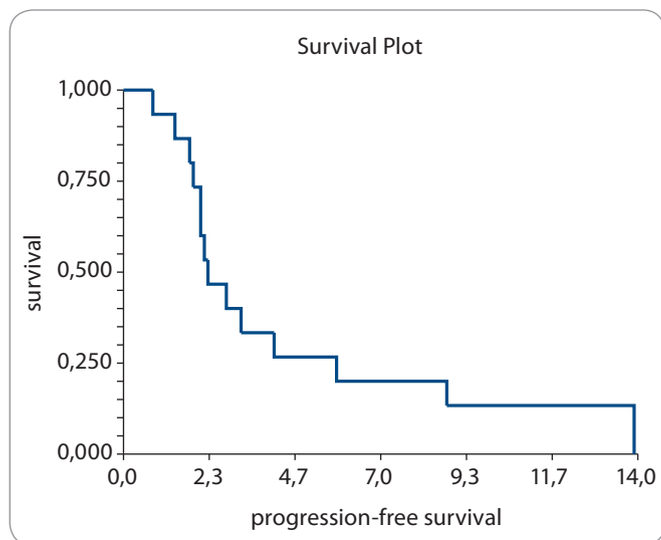
**Discussion**

Bellmunt et al introduced the first and only randomized phase III study [25] conducted in platinum-pretreated patients

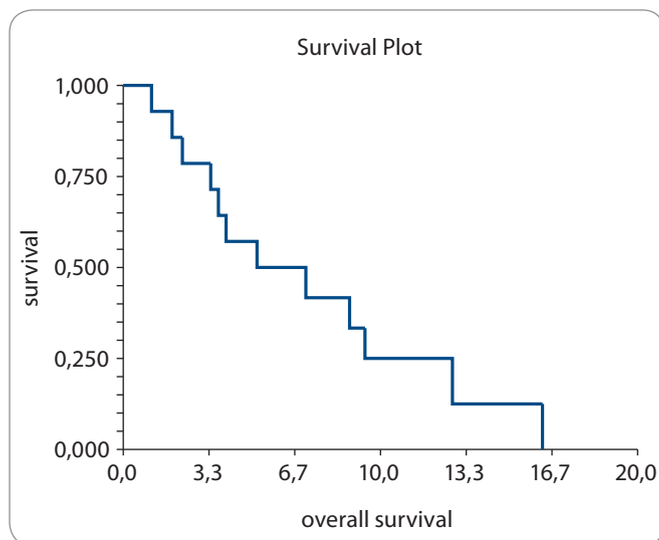
with TCCU (advanced transitional cell carcinoma of the urothelial tract) using vinflunine in the second-line setting. With reference to this study, which provided the IB level of evidence, EMEA (European Medicines Agency) has approved vinflunine in 2010. However, opponents of this study [25] argue that survival improvement of 2.6 months might be statistically significant yet, from a clinical point of view it remains irrelevant. This

might be one of the reasons why FDA (Food and Drug Administration) has never approved vinflunine as a standard second-line treatment of the advanced bladder cancer in the US. The analysis of the above-mentioned results [25], revealed several positive prognostic factors (Graph 1), such as hemoglobin level higher than 10 g/dl, the absence of liver metastases, and ECOG performance status 0–1 [27]. It is of question, whether these prognostic factors might be considered sufficient for the efficacy evaluation of a new drug like vinflunine. The analysis of 179 patients with at least one measurable target lesion showed that patients with tumor reduction ≥ 10% obtained significantly better OS than patients with tumor shrinkage lower than 10% (11,3 vs. 6,9 months, p = 0,0224). Still, even a small tumor reduction (ranging from 10%) in vinflunine treated patients led to a particular survival improvement. Hence, a decrease in SLD (sum of the longest diameter) in target lesions by about 10% might represent a promising early survival predictor in patients treated with vinflunine. SLD decreasing by ≥ 10% at the first CT control appears to be a better early outcome predictor than RECIST itself, however this remains to be confirmed by future studies.

In this prospective observational study, there were 16 patients with advanced urothelial cancer treated with second-line vinflunine in the standard



Graph 2. Progression-free survival (PFS) in the study group.



Graph 3. Overall survival (OS) in the study group.

dose. The objective response rate observed in this study was 13.33%, which is comparable to 8.60% reported by Bellmunt et al [25]. Likewise, the toxicity spectrum was similar, with neutropenia grade 3–4 being the most common side effect (37.50% in our study vs. 50.00% in Bellmunt's study), whereas the incidence of febrile neutropenia was relatively low (6.25% in this study vs. 6.0% in the study of Bellmunt et al). Constipation (grade 3–4) – the typical side effect of vinca alkaloids – was present in 18.75 patients (vs. 16,1% in Bellmunt's study). Median PFS of 2.3 months and median OS of 5.2 months obtained in this study was shorter than in the study of Bellmunt et al (3.0 months for PFS and 6.9 months for OS). This might be due to a smaller study population, as well as the pioneer nature of this prospective observational study of vinflunine treatment within Slovak boundaries.

In conclusion, single-agent vinflunine might be considered a standard of second-line treatment for patients with advanced urothelial cancer with good performance status (ECOG 0 a 1). The first Slovak experience confirms the effectivity of this treatment in terms of PFS and OS, with acceptable and well manageable toxicity profile. For patients who do not meet indication criteria, paclitaxel monotherapy could be considered with reference to a phase II study (Tab. 1). Best supportive care is optional for patients with poor performance status or when a patient refuses cytostatic treatment.

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