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# ORIGINAL RESEARCH

# Epothilones in epithelial ovarian, fallopian tube, or primary peritoneal cancer: a systematic review

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Abstract: Ovarian cancer is the most lethal gynecologic malignancy; consequently, there is a need for effective therapies. Epothilones are microtubule-stabilizing agents that inhibit cell growth. Currently, patupilone and its four synthetic derivatives ixabepilone, BMS-310705, sagopilone, 20-desmethyl-20-methylsulfanyl epothilone B and epothilone D, as well as its derivative KOS-1584, are under clinical evaluation. This is the first systematic review conducted in accordance with the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) guidelines that synthesizes all available data emerging from trials and evaluates the efficacy and safety of epothilones in epithelial ovarian, primary fallopian tube, and primary peritoneal cancer. Despite the fact that epothilones have proven active in taxane-resistant settings in preclinical models, it is not yet clear from Phase II/III studies reviewed here that their clinical activity is superior to that of taxanes. Nevertheless, responses to epothilones have been observed in platinum-refractory/resistant ovarian cancer patients. Moreover, despite the shared mechanism of action of epothilones, their clinical profile seems clearly different, with diarrhea being the most common dose-limiting toxicity encountered with patupilone, whereas neutropenia and sensory neuropathy are the most common toxic effects observed with the other epothilones. In any case, randomized trials comparing epothilones with standard treatments seem warranted to define further the role of these agents, whereas biomarker analysis might further optimize patient selection.

Keywords: ovarian cancer, epothilone, patupilone, ixabepilone, systematic review

### Introduction

Ovarian cancer is the fifth-most common cause of cancer death in women and the most lethal gynecologic malignancy.<sup>1,2</sup> In 2012, it was estimated that 22,280 women would be diagnosed with and 15,500 women die of ovarian cancer in the US.<sup>3</sup> The overall 5-year survival for 2002–2008 in 18 Surveillance, Epidemiology, and End Results geographic areas was 43.7%.<sup>3</sup> Therefore, ovarian cancer might represent an important public health issue. Landmark studies, such as GOG-111 and GOG-114, have established a platinum–taxane combination as the standard chemotherapy treatment for epithelial ovarian cancer.<sup>1,2</sup> However, more than 50% of patients with advanced disease (International Federation of Gynecology and Obstetrics stage III and IV) will relapse, requiring second-line treatment. Consequently, there is an urgent need for effective therapies for patients with relapsed ovarian cancer, particularly platinum-resistant disease.

Epothilones are microtubule-stabilizing agents that inhibit cell growth.<sup>4,5</sup> They bind to the  $\beta$ -tubulin subunit of the  $\alpha\beta$ -tubulin dimer of microtubules and induce microtubule polymerization and stabilization, resulting in G<sub>2</sub>/M arrest and the induction of apoptosis.<sup>4,5</sup> Epothilones are less susceptible than taxanes to overexpression of P-glycoprotein, the presence of certain tubulin isoforms (class III  $\beta$ -tubulin), and

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This is the first systematic review of the literature aiming to synthesize all available data emerging from trials and to evaluate the efficacy and safety of epothilones in epithelial ovarian cancer. Moreover, this systematic review deals with the administration of epothilones in primary fallopian tube and primary peritoneal cancer, given that they are managed in a similar way to epithelial ovarian cancer.

# Search strategy and data abstraction

This systematic review was performed in accordance with the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) guidelines.<sup>14</sup> The protocol of this systematic review has been submitted to the Institutional Review Board of Hippokration Hospital, Medical University of Athens, Greece, and is available upon request. Eligible articles were identified by a search of the Medline bibliographical database for the period up to September 30, 2012. The search strategy included the following keywords: ((ovarian OR ovary OR fallopian OR peritoneal OR peritoneum) AND (neoplasms OR neoplasm OR cancer OR cancers OR carcinoma OR carcinomas)) AND (epothilone OR EPO906 OR patupilone OR "BMS-247550" OR "aza-epothilone B" OR ixabepilone OR "BMS-310705" OR "KOS-862" OR "desoxyepothilone B" OR "KOS-1584" OR "ZK-EPO" OR "ZK 219477" OR sagopilone OR SH-Y03757A OR "BMS-247550").

Language restrictions were applied (only articles in English, French, and German were considered eligible); two investigators (FZ and DC), working independently, searched the literature and extracted data from each eligible study. Reviews were not eligible, while all prospective and retrospective studies, as well as case reports, were eligible for this systematic review. Manuscripts that did not state the names of the authors were excluded. In addition, we checked all the references of relevant reviews and eligible articles that our search retrieved, so as to identify potentially eligible conference abstracts.

All studies that examined the efficacy and safety of epothilones in epithelial ovarian, primary fallopian tube, or primary peritoneal cancer and reported the relevant frequencies, regardless of sample size, were considered eligible for this systematic review. For these studies, the following data were collected: first author, year of publication, agents, phase of the trial, number of patients treated, characteristics of patient population (first-line, second-line treatment, platinum-resistant, platinum-sensitive, etc), median age (years), complete response (CR) rate, partial response (PR) rate, stabilization of the disease (SD) rate, progression of the disease (PD) rate, median overall survival (OS) in months, median progression-free survival (PFS) in months, and complications. In instances where multiple (overlapping) publications stemming from the same study were identified, the larger-size study was included, unless the reported outcomes were mutually exclusive.

### Results

The search strategy retrieved 74 articles. Of these articles, 44 were irrelevant, 17 were reviews, and 13 were eligible.<sup>15-27</sup> After a search of the references of all reviews and remaining articles, 14 additional conference abstracts/articles were also included.<sup>28-41</sup> Overall, 27 studies (1,293 patients) were eligible for the systematic review (Table 1).<sup>15-39</sup> The aforementioned steps of the selection process are illustrated in detail in Figure 1.

There were seven studies evaluating patupilone (1,046 patients).<sup>15,17,21,22,24,34,35</sup> Five studies described the results of Phase I trials,<sup>17,21,22,24,35</sup> one those of a Phase II trial,<sup>34</sup> and one study involved a Phase III design.<sup>15</sup> Patupilone has been evaluated as monotherapy<sup>15,17,21,24,34,35</sup> or combined with carboplatin in epithelial ovarian, primary fallopian, or primary peritoneal cancer.<sup>22</sup> Three studies were conducted on platinum-refractory or -resistant patients,<sup>15,21,34</sup> while in one study separate data for platinum-sensitive and platinum-refractory or -resistant patients were provided;<sup>22</sup> in three studies, no data were provided regarding platinum sensitivity.<sup>17,24,35</sup> The overall response rate

	<b>Гарие Г</b> рипппагу ог ѕционеѕ еманациву цпе адплильтацой ог ероциютеѕ иг еритенагоматай, риппагу тапортал цире, али риппагу регионеат сапсет	laung une aun			spirmelial ov	arian, pr.	nary tallo	plan tup	e, aliu pi	Imary per	LONEAL CE	Incer		
Study	Chemotherapy	Patients, n	Patient	Prior	Age, years	<b>CR</b> (%)	PR (%)	SD (%)	PD (%)	Unknown	PFS,	OS,	Phase	Grade 3/4 adverse events
	regimens		characteristics	lines	(median; range)					(%)	months	months		
Colombo	Patupilone 10 mg/m <sup>2</sup>	412	Platinum-	l: 74.3%; 2:	59; 25–87		15.5* 4	43.9*	27.2*	13.3*	3.7*	13.2		Diarrhea (25.6%), nausea
et al <sup>15</sup>	over 20 minutes,	416		20.9%; ≥3:	59; 23–84	0	7.9* 4	48.3*	30.5*	13.2*	3.7*	12.7	≡	(8.2%), vomiting (8.0%),
	every 3 weeks		nt	4.9%										abdominal pain (7.7%),
	Pegylated liposomal		ovarian,	l: 73.6%; 2:										constipation (2.2%),
	doxorubicin 50 mg/m <sup>2</sup>		fallopian,	19.7%; ≥3:										abdominal distension
	every 4 weeks		or primary	6.7%										(2.7%), mucositis (0.5%),
			peritoneal											abdominal pain (upper)
			cancer (first-											(0.7%), fatigue (10.4%),
			line taxane/											pyrexia (1.0%), peripheral
			platinum-based											neuropathy (6.2%), headache
			chemotherapy)											(0.2%), decreased appetite
														(4.7%), hypokalemia (6.0%),
														dehydration (7.7%), pain in
														extremity (1.7%), back pain
														(0.2%), arthralgia (0.5%),
														anemia (4.5%), neutropenia
														(3.0%), dyspnea (4.0%),
														weight decrease (0.7%),
														insomnia (0.2%)
														Diarrhea (2.2%), nausea
														(5.9%), vomiting (5.9%),
														abdominal pain (8.6%),
														constipation (2.7%),
														abdominal distension (1.2%),
														mucositis (10.0%), abdominal
														pain (upper) (0.2%), fatigue
														(8.3%), pyrexia (1.0%),
														peripheral neuropathy
														(0.5%), decreased appetite
														(2.7%), hypokalemia (2.2%),
														dehydration (3.9%), pain in
														extremity (1.0%), back pain
														(1.0%), arthralgia (0.2%),
														anemia (3.7%), neutropenia
														(10.0%), dyspnea (3.9%), rash
														(1.7%), hand-foot syndrome
														(13.4%)
														(Continued)

Table I Summary of studies evaluating the administration of epothilones in epithelial ovarian, primary fallopian tube, and primary peritoneal cancer

Study	Chemotherapy regimens	Patients, n Patient charact	Patient Pr characteristics lin	Prior lines	Age, years CR (%) PR (%) SD (%) PD (%) Unknown PFS, (median; (%) mon range)	CR (%)	PR (%)	SD (%)	PD (%)	Unknown (%)	PFS, months	OS, months	Phase Grade 3/4 adverse events	
Tsimberidou et al <sup>17</sup>	<ul> <li>Patupilone 10 mg/m<sup>2</sup></li> <li>over 20 minutes, days 8, 29, and then every</li> <li>3 weeks (midazolam or omeprazole)</li> </ul>	6	Ovarian cancer	AA	NA		0	AN	AN	AN	ΥA	AN	_	AN
Ten Bokkel Huinink et al <sup>21</sup>		5	e d/or	One prior line with taxane, both	۲ Z	5 5		<b>N</b>	6	0.	0. E			Diarrhea (13.3%), hypokalemia (2.2%), ileus (4.4%), back pain (2.2%), pneurnoperitoneum (2.2%), perforation in anastomosis (2.2%), breast cancer (2.2%), renal insufficiency (2.2%), constipation (2.2%), ower abdominal pain (2.2%), thrombocytopenia (2.2%), subacute bowel obstruction (2.2%), dehydration (2.2%) elevation of hepatic enzymes (2.2%), dehydration (2.2%)
Forster et al <sup>22</sup>	Patupilone (dose- escalation schedule) + carboplatin 5 or 6AUC every 3 weeks	26	Platinum- sensitive ovarian or primary peritoneal cancer	l or 2	۲ Z	χ. Σ			5.4	34.5	A	ΥZ	<u>8</u>	ΔZ
Forster et al <sup>22</sup>	Patupilone (dose- escalation schedule) + carboplatin 5 or 6AUC every 3 weeks	ъ	Platinum- resistant ovarian or primary peritoneal cancer	l or 2	A	0	40.0	20.0	20.0	20.0	Ч И	AN	<u>8</u>	AN
Rubin et al <sup>24</sup>	Patupilone (dose- escalation schedule)	17	Ovarian cancer	At least I	NA	0	5.9	52.9	41.2	0	AN	٩N	_	ΝA
Smit et al <sup>34</sup>	Patupilone 10 mg/m <sup>2</sup> over 10–20 minutes, every 3 weeks	112	Platinum- refractory or -resistant ovarian, fallopian, or primary	At least 1; taxane- and platinum- pretreated	56; 18–85	0		4  .	34.8	0.71	2.5	11.2	=	Diarrhea (24%), fatigue (11%), intestinal obstruction (8%), anorexia (6%), vomiting (8%), neuropathy (4%) Grade 5: deaths (3.6%; none considered treatment-sclated)
Calvert et al <sup>35</sup>	Patupilone (dose- escalation schedule)	4		AN	NA	0	0	٩N	AN	AN	AN	AN	_	AN

Neutropenia (26.7%), peripheral neuropathy (22.2%), decreased neutrophil count (8.9%), hypokalemia (8.9%), arthralgia (8.9%), anemia (6.7%), drug hypersensitivity (6.7%), catheter- related infection (6.7%), thrombocytopenia (4.4%), farieue (4.4%), dvsnnea (4.4%)	Peripheral sensory neuropathy (8%), nausea (4%), pain in extremity (4%), vomiting (4%) Peripheral sensory neuropathy (21%), fatigue (5%), arthralgia (8%), diarrhea (3%), pain in extremity (8%), myalgia (3%), GGT increase (3%),	NA MANANA MAN	A A N	Leukopenia (14.3%), neutropenia (20.4%), anemia (8.2%), hematologic (6.1%), cardiovascular (2.0%), coagulation (4.1%), fatigue (14.3%), gastrointestinal (20.4%), gastrointestinal (20.4%), genitourinary/ renal (2.0%), hepatic (2.0%), infection (4.1%), metabolic (8.2%), neurologic (6.1%), pain (4.1%), pulmonary (4.1%) (Continued)
IS .	=	-		=
₹ Z	7.93 7.03	AN	28.4	<u>8.</u>
Υ N N	3.03	¥Ζ	A 1.4	4 4.
с. œ	26.1 17.6	0	0 X	12.2
2.8	39.1 41.2	50.0	o Z	32.7
30.6	21.7 26.5	50.0	100 NA	40.8
50.0	13.1 14.7	0	0 23.5	8.2
m œ	0 0	0	5.9	
63; 26–78	NA; 45–77 NA; 43–83	<b>∀</b> Z	A A N	۲ ۲
l: 93.3%; 2: 6.7%	l: 56%; 2: 44% l: 34%; 2: 66%	AN	NA Median 3 (range 1–6)	٩
Platinum- sensitive ovarian, fallopian, or primary peritoneal cancer	Platinum- refractory or resistant ovarian, fallopian, or primary peritoneal cancer	Fallopian and primary peritoneal cancer	n cancer m- (e (24%), m- nt (76%)	ţ
4 5	38	2		64
Sagopilone (dose- escalation schedule) + carboplatin 5AUC every 3 weeks	Sagopilone 16 mg/m <sup>2</sup> over 30 minutes every 21 days Sagopilone 16 mg/m <sup>2</sup> over 3 hours every 21 days	Sagopilone (dose- escalation schedule)	Sagopilone (dose- escalation schedule) Ixabepilone (dose- escalation schedule) + pegylated liposomal doxorubicin	Ixabepilone 20 mg/m² over I hour, days I, 8, and I5 every 28 days
McMeekin et al <sup>16</sup>	Rustin et al <sup>18</sup>	Arnold et al <sup>29</sup>	Schmid et al <sup>28</sup> Chuang et al <sup>19</sup>	De Geest et al <sup>20</sup>

				;										
	regimens		characteristics lines	lines	(median;					(%)	months	months months		
					range)									
Hensley	Gemcitabine +	œ	Ovarian cancer	e	NA	0	0	75.0	25.0	0	AN	ΑN	_	NA
et al <sup>23</sup>	ixabepilone (dose-													
	escalation schedule)													
Zhuang	Ixabepilone (dose-	6	Ovarian, primary NA	AN	NA	0	0	16.7	83.3	0	NA	ΝA	_	NA
et al <sup>25</sup>	escalation schedule)		peritoneal											
			cancer											
	Ixabepilone (dose-	10	Ovarian cancer	NA	NA	0	20.0	AN	AN	NA	AN	AN	_	NA
et al <sup>26</sup>	escalation schedule)													
Abraham	Ixabepilone (dose-	4	Ovarian cancer	NA	AN	0	0	AA	AN	AA	AA	AN	_	NA
et al <sup>27</sup>	escalation schedule)		(prior treatment											
			with paclitaxel)											
Aghajanian	Ixabepilone (dose-	2§	Ovarian, primary	3 and 4	NA	50.0	50.0	0	0	0	AN	86 and	_	NA
et al <sup>30</sup>	escalation schedule)		peritoneal	previous								l 64 days		
			cancer	lines										
Awada	Ixabepilone (dose-	18	Ovarian cancer	2 (paclitaxel/	61	0	001	0	0	0	7.1	AN	_	NA
et al <sup>31</sup>	escalation schedule)			carboplatin,										
				tamoxifen)										
Chen	lxabepilone 40 mg/m <sup>2</sup>	4	Ovarian cancer	At least 1;	AN	0	7.1	AA	AN	NA	AA	AN	=	NA
et al <sup>32</sup>	over I hour, every			taxane-										
	21 days			pretreated										
Hao et al <sup>36</sup>	Ixabepilone (dose-	3§	Ovarian cancer	AN	NA	0	0	001	0	0	AN	AN	_	NA
	escalation schedule)													
Dickson	Ixabepilone (dose-	18	Ovarian cancer	NA	NA	0	001	0	0	0	NA	AN	_	NA
et al <sup>38</sup>	escalation schedule)													
Spriggs	Epothilone D (dose-	4	Ovarian cancer	NA	NA	AA	AN	AN	AN	NA	NA	AN	_	NA
et al <sup>33</sup>	escalation schedule)													
Piro et al <sup>37</sup>	Epothilone D (dose-	AN	Ovarian cancer	NA	AN	AA	AA	AA	AA	AN	NA	AN	_	NA
	escalation schedule)													
Villalona-	KOS-1584 (dose-	2§	Ovarian cancer	NA	NA	0	0	001	0	0	NA	AN	_	NA
Calero	escalation schedule)													
et al <sup>39</sup>														
Mekhail	BMS-310705 (dose-	8	Ovarian cancer	NA	AN	0	001	0	0	0	AA	AN	_	NA
et al <sup>40</sup>	escalation schedule)													
Sessa	BMS-310705 (dose-	4	Ovarian cancer	NA	AN	0	0	AN	AN	NA	AN	AN	_	NA
et al <sup>41</sup>	escalation schedule)													

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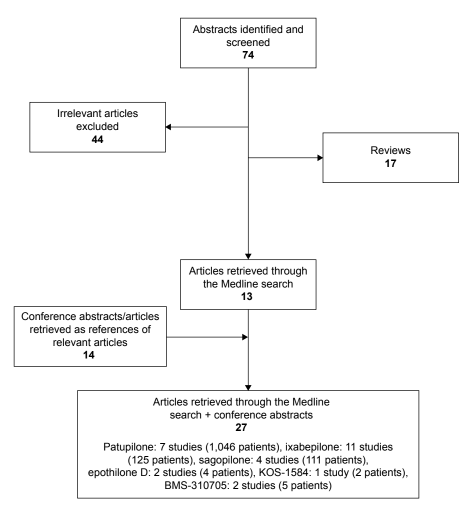


Figure I Stages of the search strategy.

(ORR) ranged between 0 and 42.3% in all studies.<sup>15,17,21,22,24,34,35</sup> Data regarding the median PFS and OS were reported in three studies:<sup>15,21,34</sup> the median PFS ranged between 2.5 and 3.7 months, while the median OS ranged between 11.2 and 14.0 months.<sup>15,21,34</sup> All studies reported detailed outcomes pertaining to efficacy and/or safety.<sup>15,17,21,22,24,34,35</sup> Further details are provided in Tables 1 and 2.

With regard to ixabepilone, eleven studies (125 patients) were identified, <sup>19,20,23,25–27,30–32,36,38</sup> nine described the results of Phase I trials, <sup>19,23,25–27,30,31,36,38</sup> and two the results of nonrandomized Phase II trials.<sup>20,32</sup> One trial dealt with platinum-refractory or -resistant epithelial ovarian cancer and primary fallopian or primary peritoneal cancer, <sup>20</sup> while in the remaining studies no detailed data were provided.<sup>19,23,25–27,30–32,36,38</sup> Moreover, ixabepilone has been evaluated as monotherapy, <sup>20,25–27,30–32,36,38</sup> as well as in combination with gemcitabine<sup>23</sup> and pegylated liposomal doxorubicin (PLD).<sup>19</sup> Clinical data regarding ixabepilone have presented promising results, with a clinical benefit rate ranging between 0 and 100%.<sup>19,20,23,25–27,30–32,36,38</sup>

Data regarding PFS and/or OS were reported in four trials (OS 28.4–14.8 months; disease-free survival 7.1–4.1 months).<sup>19,20,30,31</sup> All studies reported outcomes per-taining to efficacy and/or safety.<sup>19,20,23,25–27,30–32,36,38</sup> Further details are provided in Tables 1 and 2.

There were four studies examining sagopilone (111 patients);<sup>16,18,28,29</sup> in three of them, the agent was evaluated as monotherapy,<sup>18,28,29</sup> whereas in one sagopilone was tested in combination with carboplatin.<sup>16</sup> Two studies described the results of a Phase I trial,<sup>28,29</sup> one those of a Phase I/II trial,<sup>16</sup> and one study involved a Phase II design.<sup>18</sup> One trial was conducted in platinum-sensitive epithelial ovarian, primary fallopian, or primary peritoneal cancer,<sup>16</sup> one in platinum-refractory or -resistant cancer,<sup>18</sup> and no relevant data were provided in the other two studies.<sup>28,29</sup> ORR ranged between 0 and 58.3%, while the percentage of SD ranged between 21.7% and 100% in all studies.<sup>16,18,28,29</sup> Data regarding the median PFS and the median OS were provided in only one study.<sup>18</sup> All studies reported detailed outcomes pertaining to efficacy and/or safety (Tables 1 and 2).<sup>16,18,28,29</sup>

Parameters	Patupilone	Sagopilone	Ixabepilone	Epothilone D	KOS-1584	BMS-310705
Studies	7 (Phase I: 5; Phase II: 1;	4 (Phase I: 2;	11 (Phase I: 9;	2 (Phase I)	I (Phase I)	2 (Phase I)
	Phase III: 1)	Phase I/II: I;	Phase II: 2)			
		Phase II: I)				
Randomized trials	_	_	0	0	0	0
Patients, n	1,046	Ξ	125	4	2	5
Trials on platinum-sensitive,	3; 0; 4	1; 1; 2	1; 1; 9	0; 0; 2	0; 0; 1	0; 0; 2
platinum-resistant/refractory/						
mixed patients						
Chemotherapy agents	Patupilone (6 studies),	Sagopilone	Ixabepilone (9 studies),	Epothilone D	KOS-1584	BMS-310705
	patupilone + carboplatin	(4 studies)	ixabepilone + gemcitabine	(2 studies)	(I study)	(I study)
	(l study)		(1 study), ixabepilone +			
			pegylated liposomal			
			doxorubicin (l study)			
Main grade 3/4 adverse events						
Diarrhea (%)	13.3–25.6	0-3.0	0	NA	NA	NA
Fatigue (%)	0-11.0	0-5.0	14.3	NA	NA	AN
Neuropathy (%)	0-4.0	8.0-22.2	6.1	NA	NA	AN
Neutropenia (%)	0-3.0	0-26.7	20.4	NA	NA	AN
Nausea and vomiting (%)	2.2–16.2	0-8.0	0	NA	NA	AN
Platinum-refractory/resistant population	ation					
Median PFS (months)	2.5–3.7	2.27–3.03	4.4	NA	NA	AN
Median OS (months)	11.2–14.0	7.03-7.93	14.8	NA	NA	AN
CR (%)	0–2.8	0	6.1	NA	NA	AN
PR (%)	7.1–40	13.1–14.7	8.2	NA	NA	NA
SD (%)	20.0-43.9	21.7–26.5	40.8	NA	NA	NA
PD (%)	20.0–38.9	39.1-41.2	32.7	NA	NA	NA
Platinum-sensitive population						
Median PFS (months)	NA	NA	4.1	NA	NA	NA
Median OS (months)	NA	NA	28.4	NA	NA	AN
CR (%)	3.8	8.3	5.9	NA	NA	NA
PR (%)	38.5	50.0	23.5	NA	NA	AN
SD (%)	7.8	30.6	NA	NA	NA	NA
PD (%)	15.4	2.8	NA	٨A	AN	AN

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With regard to epothilone D, two studies with Phase I design were retrieved.<sup>33,37</sup> No data regarding CR, PR, SD, PD, PFS, OS, or detailed adverse events were reported (Tables 1 and 2). Moreover, two Phase I trials (five patients) were found, evaluating BMS-310705 in ovarian cancer patients.<sup>40,41</sup> One of these patients experienced PR, whereas detailed data regarding RRs, OS, PFS, and adverse events were not provided.<sup>40,41</sup> As far as KOS-1584 is concerned, one Phase I study (two patients) was retrieved,<sup>39</sup> showing SD in both patients with ovarian cancer. No data regarding PFS, OS, or adverse events were presented.

The characteristics of the individual studies are provided in Table 1, while the summarized features of the entire patient population are shown in Table 2. Qualitative interpretation and a critical, detailed evaluation of the individual eligible studies follow.

### Discussion

Preclinical in vitro data, stemming from a variety of taxanesensitive and -resistant cell culture models<sup>5,13</sup> together with in vivo findings in murine xenograft tumor models, have suggested that epothilones might be useful for the treatment of epithelial ovarian, primary fallopian, or primary peritoneal cancer.<sup>42,43</sup> Therefore, epothilones were evaluated in the clinical setting.

### Patupilone (epothilone B; EPO960)

Patupilone (epothilone B; EPO960) is a natural product. Based on Phase I/II data, a dose of 10 mg/m<sup>2</sup> every 3 weeks seems to be the optimal schedule for patupilone. Doselimiting toxicities include diarrhea and fatigue, whereas neurotoxicity has also been noted. Diarrhea grade 3/4 was the most common serious adverse event, ranging from 13.3% to 25.6%; however, it was manageable and usually resolved without sequelae.

Regarding the platinum-refractory/resistant population, patupilone has shown promising results, with a median PFS ranging between 2.5 and 3.7 months in heavily pretreated patients, whereas the median OS ranged between 11.2 and 14.0 months (Table 2). Moreover, the ORR ranged between 7.1% and 40.0%, whereas stabilization of the disease was achieved in 20.0%–41.1% of patients (Table 1).

A Phase I dose-escalation trial evaluated the safety, maximum tolerated dose, and pharmacokinetics of patupilone administered once every 3 weeks. In this trial, patupilone was well tolerated at doses up to 11.0 mg/m<sup>2</sup>, demonstrating promising antitumor activity (overall clinical benefit 50.1%, PFS 3 months, OS 14 months).<sup>21</sup> Adverse events were

mild to moderate in intensity, with diarrhea grade 3 (13%) being the most commonly reported serious adverse event.<sup>21</sup> The maximum tolerated dose was not reached in the study, while diarrhea, peripheral neuropathy, and fatigue were the most common dose-limiting toxicities.<sup>21</sup> The results of this trial were in accordance with those reported in a single-arm Phase II study conducted in the same patient population (platinum-refractory/resistant).<sup>34</sup> The best ORR was 7.1%, whereas 41% of patients had SD. The median PFS was 2.5 months, while the median OS was 11.2 months, with 33% of patients censored. The most common grade 3/4 toxicities were diarrhea (24%), fatigue (11%), intestinal obstruction (8%), and vomiting (8%). Finally, a study by Forster et al conducted in platinum-resistant ovarian cancer patients as a Phase IB trial (ORR 40%, SD 20%),<sup>22</sup> was in accordance with the aforementioned results. Of note, in this trial, patupilone was administered in combination with carboplatin.

Given the strong and promising clinical data, a Phase III trial was conducted evaluating patupilone in patients with resistant or refractory ovarian, fallopian, or peritoneal cancer.15 This study compared the efficacy and safety of patupilone (10 mg/m<sup>2</sup> intravenously every 3 weeks) with those of PLD (50 mg/m<sup>2</sup> intravenously every 4 weeks). There was no statistically significant difference in OS, the primary end point, between the patupilone and PLD arms (hazard ratio 0.93, 95% confidence interval [CI] 0.79–1.09; P=0.195), with median OS rates of 13.2 and 12.7 months, respectively.<sup>15</sup> Median PFS was 3.7 months for both arms. The ORR (all PRs) was higher in the patupilone arm than in the PLD arm (15.5% vs 7.9%, odds ratio 2.11, 95% CI 1.36-3.29), although disease-control rates were similar (59.5% vs 56.3%, respectively).<sup>15</sup> Frequently observed adverse events of any grade included diarrhea (85.3%) and peripheral neuropathy (39.3%) in the patupilone arm and mucositis/stomatitis (43%) and hand-foot syndrome (41.8%) in the PLD arm.15 Therefore, it seems that patupilone did not demonstrate significant improvement in OS compared with the active control, PLD.

These results led to the decision of no further development of this agent from the manufacturer. Nevertheless, a different reading of these results indicates that patupilone is active in platinum-resistant ovarian cancer, and this seems important in a chronic disease with only few active drugs being available for its platinum-resistant phase. In this context, the results of another Phase III trial evaluating patupilone vs doxorubicin in platinum-refractory/resistant ovarian, primary fallopian, or peritoneal cancer are awaited with interest (NCT00262990). As far as platinum-sensitive epithelial ovarian or primary peritoneal cancer is concerned, the only published data come from a Phase IB trial.<sup>22</sup> In this trial, patupilone was combined with carboplatin in 26 patients, showing promising results (ORR 42.3%, SD 7.8%, PD 15.4%).<sup>22</sup> It seems therefore that the role of patupilone in the treatment of platinum-sensitive epithelial ovarian cancer, if any, remains to be clarified in the future. However, studies seem warranted to evaluate its potential role in this subtype of ovarian cancer, where biomarker analysis seems mandatory to further optimize patient selection.

### Sagopilone (ZK-EPO)

Sagopilone (ZK-EPO) is a third-generation epothilone B. A dose of 16 mg/m<sup>2</sup> over 3 hours every 3 weeks seems to be the optimal schedule. With regard to toxicity, the most common adverse events are peripheral sensory neuropathy and nausea, with neuropathy grade 3 events ranging from 8.0% to 22.2% (Tables 1 and 2). Given the promising preclinical data, as well as data emerging from Phase I trials in a wide variety of cancers (also including ovarian cancer in their pools),<sup>28,29</sup> sagopilone has been evaluated in two trials on epithelial ovarian, primary fallopian, and primary peritoneal cancers.<sup>16,18</sup>

More specifically, in a Phase I/II study in women with platinum-sensitive ovarian cancer, sagopilone was tested in combination with carboplatin.<sup>16</sup> In this trial, 45 patients received sagopilone at 12 mg/m<sup>2</sup> or 16 mg/m<sup>2</sup>. The ORR was 58.3%, while the SD rate was 30.6%. The main adverse events were peripheral neuropathy (75.6%), fatigue (71.1%), and nausea (64.4%).<sup>16</sup> Therefore, it seems that sagopilone in combination with carboplatin may be effective, whereas toxicities were manageable in patients with recurrent platinum-sensitive epithelial ovarian cancer.<sup>16</sup> Moreover, the results from another randomized Phase II trial were encouraging.<sup>18</sup> In this trial, patients with platinum-refractory/resistant ovarian, primary fallopian, or peritoneal cancer were randomized to receive sagopilone 16 mg/m<sup>2</sup> as a 3- or 0.5-hourly intravenous infusion every 21 days for up to 6 weeks. The ORRs were 14.7% and 13.1%, while the median PFS was 3.03 and 2.27 months, respectively, in both arms. The 0.5-hour arm was closed when it failed to meet its target efficacy. The main drug-related adverse events were peripheral sensory neuropathy (73%; 16% grade 3), nausea (37%; 2% grade 3), fatigue (35%; 3% grade 3), and arthralgia (30%; 5% grade 3). Overall incidence of peripheral sensory neuropathy was similar in both treatment arms, with no grade 4 neuropathy events. Therefore, it seems that sagopilone may be effective, with balanced

tolerability, in patients with recurrent platinum-resistant ovarian cancer. Larger studies are nevertheless warranted to further evaluate this agent, as well as its combination with paclitaxel, in ovarian cancer patients.

# Ixabepilone (azaepothilone B; BMS-247550)

Ixabepilone (azaepothilone B; BMS-247550) is a semisynthetic analog of epothilone B. Ixabepilone has been approved by the US Food and Drug Administration in combination with capecitabine for the treatment of patients with metastatic or locally advanced breast cancer resistant to treatment with an anthracycline and a taxane, or whose cancer is taxaneresistant and for whom further anthracycline therapy is contraindicated. Moreover, it is indicated as monotherapy for the treatment of metastatic or locally advanced breast cancer in patients whose tumors are resistant or refractory to anthracyclines, taxanes, and capecitabine. The optimal schedule for ixabepilone seems to be a dose of 40 mg/m<sup>2</sup> every 3 weeks. Neutropenia is the dose-limiting toxicity, with grade 3/4 adverse events reaching 20.4% (Table 2).

Ixabepilone has been evaluated in multiple Phase I trials, where patients with ovarian cancer were included, exhibiting promising antitumor activity and a well-tolerated safety profile.<sup>19,23,25–27,30,31,36,38</sup> Of note, in a Phase I trial, ixabepilone combined with PLD was administered in 17 patients with ovarian cancer (platinum-sensitive 24%, platinum-resistant 76%).<sup>19</sup> Objective response occurred in 29% of patients (95% CI 10%–56%), median PFS was 4.1 months (95% CI 2.7–8.2 months), and median OS was 28.4 months (95% CI 7.6 months, upper limit not reached).<sup>19</sup>

Moreover, the Phase II trial, conducted by the Gynecologic Oncology Group, evaluated the efficacy and safety of ixabepilone (20 mg/m<sup>2</sup> administered over 1 hour on days 1, 8, and 15 of a 28-day cycle) in patients with recurrent or persistent platinum- and taxane-resistant primary ovarian or peritoneal carcinoma.<sup>20</sup> The ORR was 14.3% (95% CI 5.9%–27.2%).<sup>20</sup> Moreover, 40.8% of patients had SD, whereas 32.7% of them had increasing disease. The median PFS was 4.4 months (95% CI 0.8–32.6+ months), whereas the median OS was 14.8 months (95% CI 0.8–50.0).<sup>20</sup> Adverse effects included peripheral neuropathy (grade 2, 28.5%; grade 3, 6.1%), neutropenia (grade 3/4, 20.4%), fatigue (grade 3, 14.3%), and nausea (grade 3, 22%).<sup>20</sup> The results of another Phase II trial, conducted by Chen et al<sup>32</sup> were in agreement with the aforementioned.

It thus seems that ixabepilone may demonstrate antitumor activity and acceptable safety in patients with platinum- and taxane-resistant recurrent or persistent ovarian cancer. It is not clear whether this offers an advantage over retreatment with paclitaxel or docetaxel using a weekly schedule in this setting. However, larger studies of ixabepilone as monotherapy or in combination therapy in ovarian carcinoma seem warranted, with a focus on the identification of molecular markers of resistance to microtubule-stabilizing agents, including taxanes.

# Epothilone D (desoxyepothilone B; KOS-862)

Epothilone D (desoxyepothilone B; KOS-862) is a synthetic epothilone, tested in various malignancies.<sup>33</sup> Neurologic toxicity was dose-limiting in all Phase I studies of KOS-862, whereas neuropathy, fatigue, nausea, and vomiting were also observed, though to a lesser degree.<sup>8,33</sup> Epothilone D has been evaluated in two Phase I trials.<sup>33,37</sup> However, no data regarding adverse events, OS, PFS, or ORR were provided. Further trials are warranted to evaluate this agent in ovarian cancer patients.<sup>33,37</sup>

# KOS-1584 (didehydroepothilone D)

KOS-1584 (didehydroepothilone D) is a second-generation epothilone D. The most common adverse events related with the agent are gastrointestinal disorders, fatigue, and increased aminotransferase levels. KOS-1584 has shown promising antitumor activity in two patients with ovarian cancer included in a Phase I trial, as both experienced long disease stabilization.<sup>39</sup> However, the data regarding this agent in ovarian cancer are too limited to draw any firm conclusion; additional studies are needed.

# BMS-310705

BMS-310705 is an aqueous soluble, semisynthetic analog of epothilone B. The most common side effects associated with this agent are neutropenia, diarrhea, and sensory neuropathy.<sup>40,41</sup> The data on BMS-310705 in ovarian cancer patients are limited: only two Phase I trials with five patients have been published. However, one PR has been observed, showing that this agent may be active in epithelial ovarian, primary fallopian, and primary peritoneal patients. Further studies are needed to clarify the role of BMS-310705 in these patients, if any.

# Conclusion

Despite the fact that epothilones have shown activity in taxane-resistant settings in preclinical models, it is not yet clear from the Phase II/III studies reviewed here that their clinical activity is superior to that of the taxanes. Nevertheless, responses to epothilones have been observed in platinumrefractory/resistant ovarian cancer patients. Moreover, it is important to note that the availability of multiple active drugs is crucial in prolonging survival in this subpopulation of ovarian cancer, which is difficult to treat. However, it should be highlighted that patupilone is not recommended for the treatment of advanced ovarian cancer in the current clinical setting outside of an investigational trial.

Moreover, despite the shared mechanism of action of epothilones, their clinical profile is clearly different, with diarrhea being the most common dose-limiting toxicity encountered with patupilone, whereas neutropenia and sensory neuropathy are the most common toxic effects observed with the other epothilones. Additionally, combination regimens with other drugs appear feasible.

In any event, randomized trials comparing epothilones with standard treatments seem more than warranted to further define the role of these agents and/or their combinations with other existing agents, whereas biomarker analysis seems mandatory to further optimize patient selection.

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# Disclosure

The authors report no conflicts of interest in this work.

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