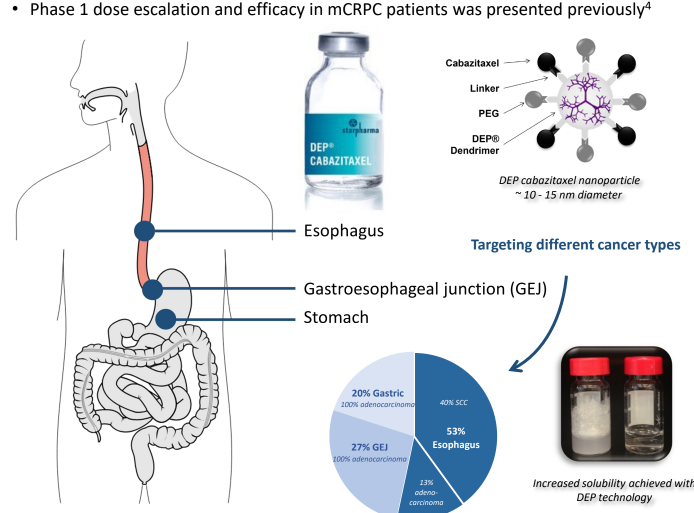
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Abstract: 374 Poster H16

Background

- Advanced esophago-gastric cancers (EGC):
- significant unmet medical need, very poor prognosis, limited available treatments - progress rapidly with 1-year survival rate of only 20%^{1,2}
- DEP cabazitaxel:
- novel, patented, poly-L-lysine dendrimer nanoparticle with polyethylene glycol (PEG), and cabazitaxel covalently attached via a hydrolysable linker
- highly water-soluble, does not contain surfactants (detergents) so does not require routine pre-medication with steroids, H2 antagonists or antihistamines, unlike conventional formulations of cabazitaxel that contain polysorbate 80
- DEP nanoparticle size restricts it to blood volume, but allows extravasation through leaky tumor vasculature³, enabling sustained delivery of cytotoxic drugs within the tumor microenvironment
- Standard cabazitaxel is widely used for treatment of metastatic castrate-resistant prostate cancer (mCRPC), but is not indicated for use in other tumor types
- Objectives of this Phase 1/2 trial were to assess preliminary efficacy and safety of DEP cabazitaxel in patients with advanced, metastatic solid cancers
- We present the efficacy and safety of DEP cabazitaxel in a cohort of locally advanced and metastatic EGC patients recruited to the Phase 2 part of this trial



Methods

• Patients with RECIST 1.1 measurable advanced EGC (squamous cell carcinoma [SCC] and adenocarcinoma) were enrolled to receive open label DEP cabazitaxel

Patients' cancer primary distribution

- DEP cabazitaxel, equivalent to 20 mg/m² cabazitaxel (recommended dose), administered intravenously (IV, ~60 min infusion), once every 21 days (3-weekly)
- Antitumor activity assessed by RECIST v1.1 and, where applicable, tumor biomarker levels such as CEA and CA 19-9; safety assessed by physical and hematological examinations, and adverse events graded according to CTCAE v4.03

KEY ELIGIBILITY CRITERIA

Inclusion Criteria	Exclusion Criteria
 Patients with advanced or metastatic solid tumors Measurable disease or evaluable tumor marker Eastern Cooperative Oncology Group (ECOG) performance status 0-1 Life expectancy ≥ 12 weeks 	 Symptomatic brain metastases or untreated spinal cord compression Absolute neutrophil count (ANC) < 1.5×10⁹/L; platelet count < 100×109/L; haemoglobin < 10 g/dL Bilirubin > ULN, or AST or ALT > 1.5 x ULN Concurrent or planned treatment with inhibitors/inducers of CYP3A4/5 Symptomatic grade 1 or ≥ grade 2 peripheral neuropathy (PN) Anti-tumor therapy ≤ 30 days or 5 half-lives prior to dosing
EU Clinical Trials Register EudraCT: 2017-003424-76	

Results

PATIENT BASELINE CHARACTERISTICS		All (N=15)	Adenocarcinoma (N=9)	Squamous Cell Carcinoma (N=6)	
Age (years)	Median (Range)	61 (25 – 73)	60 (25 – 70)	68 (50 – 73)	
Sex (% (n))	Male	67% (10)	67% (6)	67% (4)	
	Female	33% (5)	33% (3)	33% (2)	
Race (% (n))	Asian	13% (2)	11% (1)		
	Black	7% (1)	0	17% (1)	
	Black - Arabic	7% (1)	11% (1)	0	
	Caucasian	67% (10)	78% (7)	50% (3)	
	Not specified	7% (1)	0		
Prior no. lines (incl neoadjuvant / adjuvant) (% (n))	1	53% (8)	67% (6)	33% (2)	
	2	27% (4)	22% (2)	33% (2)	
	≥3	20% (3)	11% (1)	33% (2)	
Selected prior chemo-therapeutic / biologic agents (% (n))	Oxaliplatin	60% (9)	89% (8)	7% (1)	
	Cisplatin	27% (4)	11% (1)	50% (3)	
	Taxane	27% (4)	22% (2)	33% (2)	
	Anti PD-1/PD-L1	27% (4)	0	67% (4)	
Definitive Treatment (curative intent) (% (n))	Yes	47% (7)	22% (2)	83% (5)	
	FLOT + surgery	7% (1)	11% (1)	0	
	ECX + surgery	7% (1)	11% (1)	0	
	CRT	27% (4)	0	67% (4)	
	CRT + surgery	7% (1)	0	17% (1)	
	No (metastatic at diagnosis)	53% (8)	78% (7)	17% (1)	

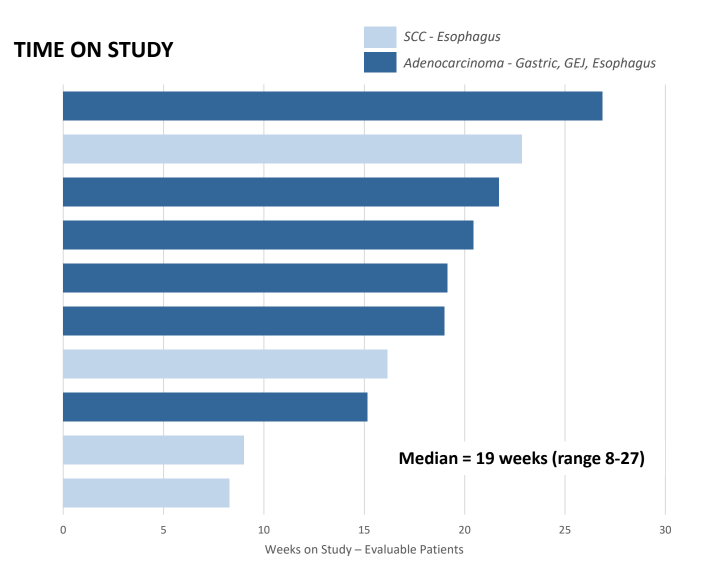
KEY OUTCOMES

- EGC patients received a median of 4.5 DEP cabazitaxel cycles (range 1-7)
- No routine steroid, antihistamine or H2 antagonist pretreatment

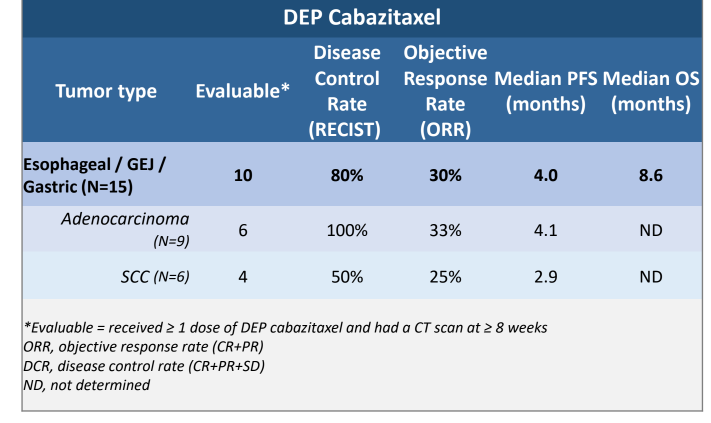
CRT = chemoradiotherapy (fluoropyrimidine / paclitaxel + platinum) + 50.4 Gy radiotherapy

CRT + surgery = carboplatin + paclitaxel + 40.4 Gy RT + esophagectomy

- Patients' prior anti-cancer therapy: median of 1 line and median of 6 cycles
- Overall ORR in evaluable patients was 30%, overall DCR was 80%
- Partial responses (PR) in both adenocarcinoma and SCC subtypes
- Stable disease (SD) for up to 27 weeks and PR for up to 17 weeks in evaluable EGC
- Increased solubility achieved with Median progression free survival (PFS) (all enrolled) was 4.0 months
 - Median Overall Survival (OS) (all enrolled) was 8.6 months



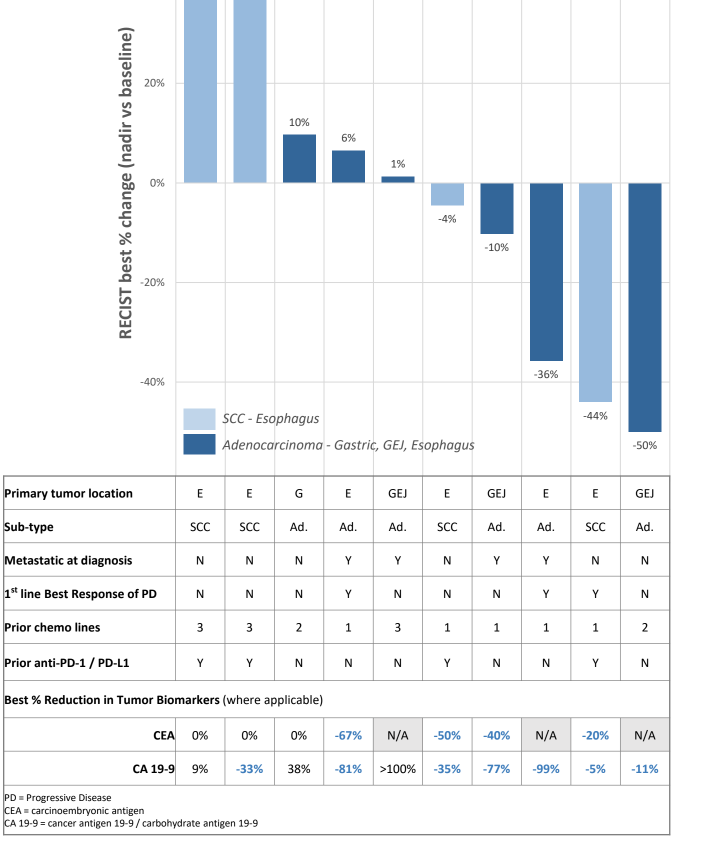
PHASE 2 ESOPHAGO-GASTRIC CANCER COHORT EFFICACY OVERVIEW



Unsolicited investigator reports of improvements in patient Quality of Life (QoL) factors

- Reduced cancer-related pain and opiate usage
- Improved performance status
- Weight gain

Best % Reduction in Tumor Target Lesion Size (RECIST 1.1)



SAFETY OVERVIEW

- DEP cabazitaxel was well-tolerated, with mostly Grade 1 (62.9%) / moderate (20%, Grade 2) treatment-related adverse events (TRAEs), with few \geq severe (grade 3)
- Limited myelosuppression, including \geq severe (grade 3) neutropenia only 1 pt had G-CSF treatment, 1 patient had secondary G-CSF prophylaxis
- Of severe TRAEs (G3/4; 17%) most (80%) were observed in 2 patients, including neutropenia, anemia, thrombocytopenia, fistula, sepsis, elevated liver enzymes

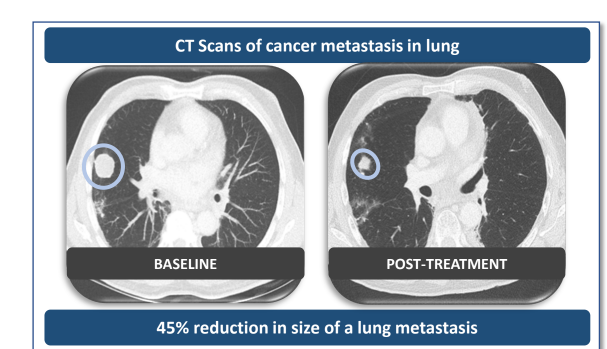
DEP Cabazitaxel Treatment-related Adverse Events (% of all TRAEs) in EGC patients

TRAEs were like those observed for standard cabazitaxel treatment

DEI Cabazitakei ilea			7,7 0.								
Grade 1	Grade 2 Grade 3		3	Grade 4							
63%	20%		7%		10%						
Treatment-Related Adverse Events (Most Extreme Grade) Experienced by ≥ 10% Patients or ≥ Grade (N=15)											
System Organ Class MedDRA Preferred Term		All Grades n (%)	Grade 1 n (%)	Grade 2 n (%)	Grade 3 n (%)	Grade 4 n (%)					
Blood and Lymphatic System D	isorders										
	Anemia	7 (47)	1 (7)	4 (27)	2 (13)						
	Neutropenia	7 (47)	2 (13)	1 (7)	1 (7)	3 (20)					
Thror	nbocytopenia	2 (13)	1 (7)		1 (7)						
Gastrointestinal Disorders											
Esophago-puln	nonary fistula	1 (7)				1 (7)					
	Nausea	5 (33)	5 (33)								
	Vomiting	3 (20)	2 (13)	1 (7)							
General Disorders and Adminis	tration Site C	onditions									
	Fatigue	5 (33)	4 (27)	1 (7)							
Infections and infestations											
	Sepsis	1 (7)				1 (7)					
Investigations											
Alanine aminotransferase (A	ALT) increased	2 (13)	1 (7)			1 (7)					
Aspartate aminotransferase (A	ST) increased	3 (20)	2 (13)			1 (7)					
White blood cell cou	ınt decreased	2 (13)		1 (7)	1 (7)						
Nervous system Disorders											
Periphero	al neuropathy	9 (60)	5 (33)	3 (20)	1 (7)						
Skin and Subcutaneous Tissue	Disorders										
	Nail distrophy	2 (13)	2 (13)								

CASE REPORT: 73-year-old man with Stage IV esophageal SCC

- Developed multiple new lung metastases within 3 months of completing an initial course of chemoradiotherapy with curative intent (first line)
- Achieved partial response following 5 cycles of DEP cabazitaxel:
 - 44% overall decrease in tumor burden
 - 45% reduction in size of a lung metastasis
 - Disease control maintained for 24 weeks



DEP CABAZITAXEL MONOTHERAPY (Q3W DOSE REGIMEN)

HIGHLY ENCOURAGING ANTI-TUMOR ACTIVITY IN ADVANCED ESOPHAGO-GASTRIC CANCERS OF BOTH ADENOCARCINOMA AND SCC HISTOLOGY

- durable responses in esophageal, GEJ and gastric cancers
- concomitant tumor biomarker reductions

WELL-TOLERATED IN ADVANCED CANCER PATIENTS WITH HIGH-RISK **CLINICAL FEATURES INCLUDING POOR NUTRITIONAL STATUS**

- mostly mild / moderate TRAEs
- limited myelosuppression, lack of need for G-CSF treatment or prophylaxis
- no severe hypersensitivity even in the absence of routine steroid, H2 antagonist and antihistamine premedication

Conclusion

DEP cabazitaxel administered as a monotherapy, once every 3 weeks exhibited highly encouraging anti-tumor activity in >1L, advanced EG cancers:

- multiple anatomic locations (esophagus, GEJ and stomach)
- different histological sub-types: adenocarcinoma and SCC
- included many patients who were refractory to 1st line therapy
- achieved excellent efficacy responses:
 - median progression-free survival (PFS) of 4.0 months
 - median overall survival (OS) of 8.6 months
 - 30% ORR and 80% DCR in evaluable patients

The DEP cabazitaxel results compare very favorably to standard-of-care paclitaxel treatment in patients with esophageal or gastro-esophageal junction cancers, with DEP cabazitaxel achieving a more than 50% longer median PFS and a 29% longer median OS than paclitaxel administered weekly as a second-line treatment⁵. The DEP cabazitaxel efficacy results in advanced EGC patients, along with highly encouraging efficacy results in patients with mCRPC and platinum-resistant ovarian cancer, indicate the promising clinical potential of DEP cabazitaxel in a range of cancer types, including cancers for which conventional cabazitaxel is not indicated⁶.

ACKNOWLEDGEMENTS

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Study sponsored by Starpharma Pty Ltd, Abbotsford, Australia

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