CLINICAL TRIALS

Open clinical uro-oncology trials in Canada

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BLADDER CANCER

A MULTICENTRE, RANDOMIZED PLACEBO-CONTROLLED, DOUBLE-BLIND PHASE III TRIAL OF SINGLE-DOSE INTRAVESICAL EOQUIN (APAZIQUONE) AS A SURGICAL ADJUVANT INSTILLED IN THE EARLY POST-OPERATIVE PERIOD IN PATIENTS UNDERGOING TRANSURETRHAL RESECTION FOR NONINVASIVE BLADDER CANCER

Trial ID:	SPI-612	
Coordination:	Spectrum Pharmaceuticals	
Trial design:	Phase III, blinded.	
Patient population:	Patients with resected bladder carcinoma TA, G1/G2.	
Sample size		
& primary endpoint:	n = 674, local recurrence at 2 years	
RANDOMIZED STUDY OF	FLAROTAXEL + CISPLATIN (LC) VS. GEMCITABINE + CISPLATIN (GC) IN THE FIRST	
LINE TREATMENT OF LO	CALLY ADVANCED/METASTATIC UROTHELIAL TRACT OR BLADDER CANCER	
Trial ID:	NCT00625664, EFC6668	
Coordination:	sanofi-aventis	
Trial design:	Randomized, open-label, multi-center study comparing the efficacy and safety of	
	XRP9881 plus cisplatin to gemcitabine plus cisplatin.	
Patient population:	First line treatment of locally advanced/metastatic urothelial tract or bladder cancer.	
Sample size		
& primary endpoint:	n = 900, overall survival	
A RANDOMIZED, PLACE	BO-CONTROLLED PHASE II STUDY TO COMPARE THE EFFICACY AND SAFETY	
OF SU011248 PLUS BEST	SUPPORTIVE CARE (BSC) VERSUS PLACEBO PLUS BSC IN PATIENTS WITH	
ADVANCED UROTHELIA	L TRANSITIONAL CELL CARCINOMA WHO HAVE FAILED OR ARE INTOLERANT	
TO CISPLATIN CONTAIN	ING CHEMOTHERAPY	
Trial ID:	SPRUCE	
Coordination:	Canadian Urologic Oncology Group (CUOG)	
Trial design:	A randomized phase II study comparing sunitinib to placebo.	
Patient population:	Recurrent or metastatic transitional cell carcinoma failed, intolerant of, or ineligible	
	for first-line cisplatin-based combination chemotherapy.	

Sample size	*
& primary endpoint:	n = 58, progression-free survival

PROSTATE ADENOCARCINOMA

LOCALIZED PROSTATE CANCER

Low Risk

A PHASE III STUDY OF ACTIVE SURVEILLANCE THERAPY AGAINST RADICAL TREATMENT IN PATIENTS		
DIAGNOSED WITH FAVORABLE RISK PROSTATE CANCER (START)		
Trial ID:	NCIC CTG PR11	
Coordination:	National Cancer Institute of Canada Clinical Trials Group (NCIC CTG)	
Trial design:	A phase III study comparing radical prostatectomy or radical radiotherapy at the	
Patient population:	time of initial diagnosis to active surveillance and selective intervention based on pre-specified biochemical, histological or clinical criteria. Suitable candidates for radical prostatectomy or radiotherapy. No previous treatment for prostate cancer for greater than 6 months. Favorable risk as defined by the following:	
	clinical stage T1b, T1c, T2a or T2b, surgical Gleason score <= 6, PSA <= 10.0 ng/ml.	
Sample size		
& primary endpoint:	n = 2130, disease specific survival	

Intermediate Risk

A PHASE III PROSPECTIVE RANDOMIZED TRIAL OF DOSE-ESCALATED RADIOTHERAPY WITH OR WITHOUT SHORT TERM ANDROGEN DEPRIVATION THERAPY FOR PATIENTS WITH INTERMEDIATE RISK PROSTATE CANCER

Trial ID:	RTOG 0815	
Coordination:	Radiation Therapy Oncology Group (RTOG)	
Trial design:	A randomized controlled trial to demonstrate an overall survival (OS) advantage for the addition of short term (6 months) ADT versus no additional ADT in the context of dose escalated RT for patients with intermediate risk prostate cancer.	
Sample size		
& primary endpoint:	n = 1520, overall survival	
PROSTATE FRACTIONAT	ED IRRADIATION TRIAL (PROFIT)	
Coordination:	Ontario Clinical Oncology Group (OCOG)	
Trial design:	A phase III study assessing the relative efficacy of dose-escalated radiation therapy (78 Gy	
	in 39 fractions) versus a hypofractionated course of radiation (6000 Gy in 20 fractions).	
Patient population:	Intermediate-risk prostate cancer.	
Sample size		
& primary endpoint:	n = 1204, biochemical (PSA) failure	

High Risk

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RANDOMIZED PHASE III STUDY OF NEO-ADJUVANT DOCETAXEL AND ANDROGEN DEPRIVATION		
PRIOR TO RADICAL PROSTATECTOMY VERSUS IMMEDIATE RADICAL PROSTATECTOMY IN PATIENTS		
WITH HIGH-RISK, CLIN	ICALLY LOCALIZED PROSTATE CANCER	
Trial ID:	NCIC PRC3	
Coordination:	Intergroup (Cancer and Leukemia Group B)	
Trial design:	A phase III comparison of neoadjuvant chemohormonal therapy with goserelin or	
	leuprolide for 18-24 weeks with docetaxel IV every 3 weeks for up to six courses	
	followed by radical prostatectomy with staging pelvic lymphadenectomy versus	
	radical prostatectomy with staging lymphadenectomy alone.	
Patient population:	High-risk prostate cancer.	
Sample size		
& primary endpoint:	n = 750, 3 year biochemical progression-free survival	

POST-RADICAL PROSTATECTOMY

RADICALS: RADIOTHERAPY AND ANDROGEN DEPRIVATION IN COMBINATION AFTER LOCAL SURGERY		
Trial ID:	NCIC PR13	
Coordination:	Intergroup (MRC)	
Trial design:	Phase III clinical trial with randomizations both for radiotherapy timing, and for	
	hormone treatment duration.	
Patient population:	Men who have undergone radical prostatectomy for prostatic adenocarcinoma within	
	3 months, post-operative serum PSA less than 0.4 ng/ml. Uncertainty in the opinion	
	of the physician and patient regarding the need for immediate post-operative RT.	
Sample size		
& primary endpoint:	n = 5100, disease free survival	

BIOCHEMICALLY RELAPSED PROSTATE CANCER

A MULTICENTER CLINIC RECURRENT PROSTATE Trial ID: Coordination: Trial design: Patient population: Sample size & primary endpoint:	CAL STUDY OF THE SONABLATE® 500 (SB-500) FOR THE TREATMENT OF LOCALLY CANCER WITH HIFU FSI-003 Focus Surgery Inc. Single arm phase II. Men with locally recurrent prostate cancer following external beam irradiation. n = 202, absence of biochemical failure and negative prostate biopsy rate at 12 months
	ARISON OF IMMEDIATE VERSUS DEFERRED ANDROGEN DEPRIVATION THERAPY RECURRENT PROSTATE CANCER AFTER RADICAL RADIOTHERAPY ELAAT OCOG A phase III trial comparing immediate to deferred androgen deprivation therapy. Patients who have undergone prior radical radiation for prostate cancer and are now experiencing a biochemical recurrence.
Sample size & primary endpoint:	n = 1100, time to androgen independent disease
A RANDOMIZED, DOUBLE-BLIND, MULTICENTRE PHASE II CONTROLLED TRIAL ASSESSING ZACTIMA (VANDETANIB) AGAINST PLACEBO IN PROLONGING THE OFF-TREATMENT INTERVAL IN PROSTATE CANCER SUBJECTS UNDERGOING INTERMITTENT ANDROGEN DEPRIVATION HORMONAL THERAPY Trial ID:Trial ID:ZENITH/D4200L00010Coordination:CURC/CUOGTrial design:Randomized double-blind placebo-controlled phase II.Patient population:Men with rising PSA after treatment for localized prostate cancer.Sample sizen = 100, PSA > 5 ng/l by 52 weeks in the off treatment interval during intermittent androgen ablation therapy.	
	ORT-TERM ANDROGEN DEPRIVATION WITH PELVIC LYMPH NODE OR PROSTATE RAPY (SPPORT) IN PROSTATE CANCER PATIENTS WITH A RISING PSA AFTER OMY RTOG 0534 RTOG

Irial ID:	RTOG 0534
Coordination:	RTOG
Trial design:	Phase II comparing radiotherapy alone to radiotherapy with short-term androgen deprivation.
Patient population:	Males who have undergone radical prostatectomy, followed by PSA rise to > 0.2 ng/ml.
Sample size	
& primary endpoint:	n = 1764, 5-year freedom from progression

A STUDY OF ANDROGEN DEPRIVATION WITH LEUPROLIDE, +/- DOCETAXEL FOR CLINICALLY ASYMPTOMATIC PROSTATE CANCER SUBJECTS WITH A RISING PSA

XRP6976J/3503
sanofi-aventis
A phase III comparison of and rogen deprivation with or without docetaxel in men with rising
PSA followed by radical prostatectomy.
No metastases and PSA doubling time ≤ 9 months
n = 412, progression-free survival

METASTATIC PROSTATE CANCER

A RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED PHASE III STUDY OF EARLY VERSUS STANDARD ZOLEDRONIC ACID TO PREVENT SKELETAL RELATED EVENTS IN MEN WITH PROSTATE CANCER METASTATIC TO BONE

Trial ID:	NCIC PRC2	
Coordination:	Intergroup (Cancer and Leukemia Group B)	
Trial design:	A phase III study comparing treatment with zoledronic acid at the time of initiation of	
	androgen deprivation therapy for metastatic prostate cancer to treatment at time of progression to hormone-refractory disease.	
Patient population:	Metastatic prostate cancer with at least one bone metastasis by radiographic imaging receiving androgen deprivation therapy.	
Sample size & primary endpoint:	n = 680, time to first skeletal related event	

CASTRATE RESISTANT PROSTATE CANCER

A PHASE III TRIAL OF ZD4054 (ENDOTHELIN A ANTAGONIST) IN NON-METASTATIC HORMONE RESISTANT PROSTATE CANCER

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Trial ID:	ENTHUSE M0/D4320C00015
Coordination:	AstraZeneca
Trial design:	Placebo controlled phase III randomized
Patient population:	HRPC with rising PSA after surgical or medical castration but no evidence of
	metastases.
Sample size	

& primary endpoint:	n = 1500, progression-free survival
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A PHASE III, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED STUDY OF ABIRATERONE ACETATE (CB7630) PLUS PREDNISONE IN ASYMPTOMATIC OR MILDLY SYMPTOMATIC PATIENTS WITH METASTATIC CASTRATION RESISTANT PROSTATE CANCER Trial ID: COU-A A-302

IIIal ID.	COO-AA-302
Coordination:	Cougar Biotechnology, Inc.
Trial design:	Randomized double-blind placebo-controlled phase III
Patient population:	Men with minimally symptomatic metastatic castration resistant prostate cancer and
	no prior cytotoxic chemotherapy.
Sample size	

& primary endpoint: n = 1000, overall survival and progression-free survival

A PHASE III, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED STUDY TO ASSESS THE EFFICACY AND SAFETY OF 10 MG ZD4054 IN COMBINATION WITH DOCETAXEL IN COMPARISON WITH DOCETAXEL IN PATIENTS WITH METASTATIC HORMONE-RESISTANT PROSTATE CANCER

Trial ID:	ENTHUSE M1C/D4320C00033
Coordination:	AstraZeneca
Trial design:	Placebo controlled phase III trial
Patient population:	Metastatic HRPC
Sample size	
& primary endpoint:	n = 1044, overall survival

AFLIBERCEPT VERSUS PI	DOMIZED, DOUBLE-BLIND STUDY COMPARING THE EFFICACY AND SAFETY OF LACEBO EVERY 3 WEEKS IN PATIENTS TREATED WITH DOCETAXEL/PREDNISONE ROGEN INDEPENDENT PROSTATE CANCER VENICE/EFC6546 sanofi-aventis A phase III study comparing the addition of aflibercept to standard docetaxel/ prednisone. Metastatic hormone-refractory prostate cancer and no prior palliative chemotherapy.	
& primary endpoint:	n = 1200, overall survival	
A PHASE II STUDY OF S CANCER AFTER FIRST L Trial ID: Coordination: Trial design: Patient population: Sample size & primary endpoint:	SU011248 FOR MAINTENACE THERAPY IN HORMONE REFRACTORY PROSTATE INE CHEMOTHERAPY SMART/TBCC-0707001 Tom Baker Cancer Centre Phase II. Patients with HRPC in remission after docetaxel. n = 30, progression-free survival	
SAFETY STUDY OF ORAL CANCER PREVIOUSLY T Trial ID: Coordination: Trial design: Patient population:	ASE III, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED EFFICACY AND MDV3100 IN PATIENTS WITH PROGRESSIVE CASTRATION-RESISTANT PROSTATE REATED WITH DOCETAXEL-BASED CHEMOTHERAPY AFFIRM ProTrials Research Inc./Medivation Inc. Randomized (2:1), double-blind, multicenter study comparing MDV3100 to placebo. Metastatic castration-resistant prostate cancer progressive despite prior docetaxel or mitoxantrone chemotherapy.	
Sample size	n – 1170. ovorall survival	
& primary endpoint:n = 1170, overall survivalA DOUBLE-BLIND, RANDOMIZED, MULTIPLE DOSE, PHASE III, MULTICENTER STUDY OF ALPHARADIN IN THE TREATMENT OF PATIENTS WITH SYMPTOMATIC HORMONE REFRACTORY PROSTATE CANCER WITH SKELETAL METASTASESTrial ID:ALSYMPCACoordination:Algeta ASATrial design:Randomized, double-blind, multicenter study comparing Alpharadin (radium-223) to placebo.Patient population:Metastatic castration-resistant prostate cancer progressive despite prior docetaxel or mitoxantrone chemotherapy.Sample size & primary endpoint:n = 750, overall survival		
& primary endpoint:	n = 750, overall survival	

RENAL CELL CANCER

A RANDOMIZED, DOUBLE-BLIND PHASE III TRIAL OF ADJUVANT SUNITINIB VERSUS SORAFENIB VERSUS		
PLACEBO IN PATIENTS WITH RESECTED RENAL CELL CARCINOMA (ASSURE)		
Trial ID:	NCIC REC.2	
Coordination:	Intergroup (ECOG)	
Trial design:	A phase III surgical adjuvant study assessing the effectiveness of sunitinib or sorafenib compared to placebo.	
Patient population:	Resected renal cell carcinoma, T1b grade 3-4 or higher and/or N+.	
Sample size		
& primary endpoint:	n = 1332, overall survival	
A STUDY OF PAZOPANIB VERSUS SUNITINIB IN THE TREATMENT OF SUBJECTS WITH LOCALLY ADVANCED AND/OR METASTATIC RENAL CELL CARCINOMA		
Trial ID:	COMPARZ/VEG108844	
Coordination:	GlaxoSmithKline	
Trial design:	A phase III study comparing pazopanib to sunitinib in metastatic renal carcinoma.	
Patient population:	Untreated metastatic clear cell renal carcinoma.	
Sample size		
& primary endpoint:	n = 876, progression-free survival	
A RANDOMIZED TRIAL OF TEMSIROLIMUS AND SORAFENIB AS SECOND LINE THERAPY IN PATIENTS WITH ADVANCED RENAL CELL CARCINOMA WHO HAVE FAILED FIRST LINE SUNITINIB THERAPY		
Trial ID:	3066K1-404-WW	
Coordination:	Wyeth	
Trial design:	An international, randomized, open label, multicenter phase III study assessing weekly temsirolimus versus sorafenib twice daily in the second line setting.	
Patient population:	Histologically confirmed metastatic renal cell carcinoma, progressive disease on sunitinib.	
Sample size		
& primary endpoint:	n = 440, progression-free survival and safety	