

**REFINING RISK-STRATIFICATION OF HIGH-RISK AND LOCOREGIONAL  
PROSTATE CANCER: A POOLED ANALYSIS OF RANDOMIZED TRIALS**

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## 1    **ABSTRACT**

2    **Background:** Radiotherapy (RT) and long-term ADT (ltADT; 18-36 months) is a  
3    standard-of-care in the treatment of high-risk localized/locoregional prostate cancer  
4    (HRLPC). We evaluated outcomes in patients treated with RT + ltADT to identify which  
5    patients have poorer prognosis with standard therapy.

6    **Methods:** Individual patient data (IPD) from patients with HRLPC (as defined by any of  
7    the following 3 risk factors [RFs] in context of cN0 disease: Gleason score  $\geq 8$ , cT3-T4,  
8    PSA  $>20\text{ng/mL}$ , or cN1) treated with RT and ltADT on randomized controlled trials  
9    collated by the Intermediate Clinical Endpoints in Cancer of the Prostate group.  
10   Outcome measures of interest were metastasis-free survival (MFS), overall survival  
11   (OS), time to metastasis (TTM) and prostate cancer-specific mortality (PCSM).  
12   Multivariable Cox and Fine-Gray regression estimated hazard ratios (HR) for the 3 RFs  
13   and cN1 disease.

14   **Findings:** 3604 patients from 10 trials were evaluated, with a median PSA of  $24\text{ng/mL}$ .  
15   Gleason score  $\geq 8$  (MFS HR=1.45; OS HR=1.42), cN1 disease (MFS HR=1.86; OS  
16   HR=1.77), cT3-4 disease (MFS: HR=1.28; OS: HR=1.22), and PSA  $>20\text{ng/mL}$  (MFS  
17   HR=1.30; OS HR=1.21) were associated with poorer outcomes. Adjusted 5-year MFS  
18   rates were 83% and 78% for patients with 1 and 2-3 RFs, and 10-year MFS rates were  
19   63% and 53%, respectively; corresponding 10-year adjusted OS rates were 67% and  
20   60%. In cN1 patients, adjusted 5- and 10-year MFS rates were 67% and 36%,  
21   respectively, and 10-year OS was 47%.

**Conclusion:** HRLPC patients with 2-3 RFs (and cN0) or cN1 disease had the poorest outcomes on RT and ItADT. This will help in counselling patients treated in routine practice and in guiding adjuvant trials in HRLPC.

## 1 INTRODUCTION

2 Approximately 25% of localized prostate cancers are considered 'high-risk', as defined  
3 by a Gleason score  $\geq 8$  and/or PSA  $> 20$  ng/mL and/or clinical T3/T4 disease,[1] with  
4 evidence of regional nodal involvement seen in an additional 10-15% of cancers.[2]  
5 Together, high-risk and locoregional prostate cancer (HRLPC) are associated with a  
6 significant risk of prostate cancer mortality and account for two-thirds of deaths from  
7 prostate cancer at 10 years.[3]

8 Multimodal therapy is usually required for HRLPC, with RT and long-term (lt; 18-36  
9 months) androgen deprivation therapy (ADT) being a widely accepted standard-of-  
10 care.[4, 5] Recently, the STAMPEDE trial showed a significant improvement in  
11 metastasis-free (MFS) and overall survival (OS) with the addition of abiraterone to RT  
12 and ltADT in men with HRLPC, as defined by either cN1 disease or two of: Gleason  $\geq 8$ ,  
13 cT3-4 and PSA  $\geq 40$  ng/mL.[6] The STAMPEDE participants represented a particularly  
14 high-risk group, with a median PSA of 30-40 ng/mL and 40% of patients having N1  
15 disease on conventional imaging. Trials evaluating other novel androgen receptor  
16 pathway inhibitors (ARPIs) in combination with RT and ADT for HRLPC are ongoing and  
17 are being powered with the assumption of 5-year MFS of ~75% in the control arm of RT  
18 + ltADT. (**Supplementary Table 1**).

19 Based on these considerations, we sought to evaluate long-term outcomes in various  
20 groups of patients with HRLPC treated with RT and ltADT on randomized trials, whose  
21 individual patient data (IPD) are available within the Intermediate Clinical Endpoints in  
22 Cancer of the Prostate (ICECaP) data repository.[7] Specifically, we aimed to define the



outcomes for a range of endpoints – including MFS and OS, but also cancer-specific measures such as time to metastasis (TTM) and prostate cancer-specific mortality (PCSM) – associated with different permutations of standard clinicopathological variables. Defining the patients with HRLPC with the poorest outcomes may help clarify those most likely to benefit from treatment intensification as well as those who may achieve excellent outcomes with RT and ItADT alone and be candidates for treatment de-intensification.

## METHODS

### *Trial and Patient Selection*

The ICECaP repository comprises trials collected in the initial meta-analysis that has been previously published[8] as well as data from additional trials collected between May 2020 and February 2023 since this publication; the meta-analysis was conducted with adherence to PRISMA guidelines. For the current study, only IPD from patients in RT-based trials who had HRLPC and were treated with 18-36 months of ADT were eligible; HRLPC was defined as cN1 disease (on conventional imaging) and/or any of Gleason  $\geq 8$ , cT3-4 and PSA  $> 20$  ng/mL. A flowchart of selection of patients for this study is shown in **Supplementary Figure 1**, and the list of eligible patients from included trials is provided in **Supplementary Table 2**.

### *Definition of endpoints*

The clinical outcomes analyzed were MFS, OS, TTM and PCSM. MFS was measured from the date of randomization to date of first evidence of distant metastases (by conventional imaging – CT, MRI and/or bone scan – or histology) or death from any cause; or censored at the date of most recent follow-up. TTM was defined analogously to MFS but non-prostate cancer deaths without prior disease progression were counted as a competing risk. OS was measured from the date of randomization to death from any cause, or censored at the date of most recent follow-up in patients who were alive. PCSM was defined similarly as OS, but non-prostate deaths were considered as a competing risk.

1

## 2 *Statistical Analysis*

3 5-year MFS and OS were estimated by the Kaplan Meier method; 5-year of TTM and  
4 PCSM were estimated using cumulative incidence function accounting for competing risk.  
5 Multivariable Cox regression models (for MFS and OS) and the Fine and Gray Competing  
6 risks regression (for TTM and PCSM) were performed to estimate the strength of  
7 association of clinical outcomes with pre-defined baseline risk factors, including biopsy  
8 Gleason ( $\geq 8$  vs.  $\leq 7$ ), clinical T-stage (cT3-4 vs. cTx1-2), PSA at randomization ( $< 10$ ng/mL,  
9  $10$ - $20$ ng/mL, and  $> 20$ ng/mL), and clinical N-stage (cN1 vs. cN0). The PSA cutoffs were  
10 based on established risk stratification criteria for localized prostate cancer.[9] These  
11 models were adjusted for age at randomization, ADT duration ( $\geq 24$  months vs. 18 months)  
12 and radiotherapy dose ( $\leq 70$ Gy,  $> 70$ Gy and unknown) and stratified by years of enrollment  
13 (per 5-year increment) to account for variability of follow up times across the trials. Median  
14 follow-up was calculated using the reverse Kaplan-Meier method.

15 Based on number of baseline adverse risk factors from the multivariable models above,  
16 we estimated adjusted 5- and 10-year MFS and OS from Cox regression[10] and adjusted  
17 5- and 10-year TTM and PCSM[11] from Fine and Gray regression models. Additionally,  
18 we reported unadjusted Kaplan Meier estimates of MFS and OS and unadjusted  
19 cumulative incidence of TTM and PCSM for various pre-planned risk subgroups (by  
20 permutations of Gleason, clinical T-stage, PSA and clinical N stage) as well as for post-  
21 hoc analyses of number of adverse factors by age ( $\leq$  vs  $> 68$  years) and radiotherapy dose  
22 delivered ( $\leq 70$ Gy,  $> 70$ Gy and unknown) using the median as a threshold for each

1 stratification variable. The adjusted survival curves were estimated using R  
2 “adjustedCurve” package (<https://www.r-project.org/>). All other statistical analyses were  
3 performed using the SAS software application (version 9.4; SAS Institute, Cary, NC,  
4 USA). Two-sided p values <0.05 were considered statistically significant.

## RESULTS

A total of 3604 patients with HRLPC treated across 10 trials evaluating RT and ItADT were eligible. Baseline characteristics of these patients at the time of randomization are shown in **Table 1**. Median age was 68 years and median PSA was 24ng/mL; 1942 patients (54%) had Gleason 8-10 disease, 2061 (57%) had a PSA >20ng/mL, 2602 (72%) were cT3-4, and 422 (12%) had cN1 disease. Median follow-up was 8.6 years (interquartile range 6.0-11.8), and 5-year MFS and OS rates in the entire population were 78% (95% CI 77-80) and 84% (83-85), respectively.

**Table 2** shows the results of multivariable analyses evaluating the adjusted associations of clinical risk factors with long-term outcomes. Statistically significant associations were seen for Gleason score  $\geq 8$  (MFS HR=1.45 [95% CI 1.29-1.63]; OS HR=1.42 [1.26-1.61]), cN1 disease (MFS HR=1.86 [1.56-2.21]; OS HR=1.77 [1.45-2.15]), cT3-4 disease (MFS HR=1.28 [1.13-1.45]; OS HR=1.22 [1.07-1.39]), and PSA >20ng/mL (MFS HR=1.30 [1.13-1.50]; OS HR=1.21 [1.05-1.41]). Broadly similar trends were seen in the associations between these variables and TTM and PCSM.

Given the variability in associations between the clinicopathological variables and outcomes, we generated Kaplan-Meier estimates of 5- and 10-year MFS rates based on various permutations of risk factors (Gleason 7 vs  $\geq 8$ , PSA <10 vs 10-20 vs  $\geq 20$ ng/mL, cT3-4 vs cTx1-2, cN1; **Table 3**); estimates of 5- and 10-year OS, TTM and PCSM are shown in **Supplementary Table 3**. Overall, outcomes were best in cN0 patients with just one adverse risk factor (Gleason  $\geq 8$ , PSA >20ng/mL, cT3-4), intermediate in patients with 2 adverse risk factors and worse in patients with all 3 risk factors; the

1 poorest outcomes overall were seen in patients with cN1 disease regardless of other  
2 risk factors.

3 Given the similar outcomes between cN0 patients with 2 or 3 adverse risk factors, these  
4 were grouped together and adjusted survival curves showing MFS and OS, and  
5 cumulative incidence of TTM and PCSM based on number of risk factors (1 vs. 2-3 vs.  
6 cN1) are shown in **Figure 1**. Adjusted 5- and 10-year estimates of MFS, OS, TTM and  
7 PCSM rates by these risk groups (1 vs. 2-3 vs. cN1) are shown in **Table 4**. Adjusted 5-  
8 year MFS rates were 83% (81-85), 78% (76-79) and 67% (62-71) for patients with 1, 2-3  
9 risk factors and cN1 disease, respectively, while corresponding adjusted 5-year OS  
10 rates were 87% (86-88), 84% (82-85) and 77% (74-80). Similar trends in outcomes by  
11 risk groups were seen when stratifying by age or RT dose (**Supplementary Tables 4-**  
12 **5**), with generally better outcomes seen across risk groups in patients treated at higher  
13 RT doses.

14 We also evaluated the STAMPEDE definition of high-risk in our cohort (i.e. cN1 or  
15 Gleason 8-10, cT3-4, PSA  $\geq$ 40ng/mL), which led to a decrease in the number of  
16 patients with 2-3 risk factors. Despite the higher PSA cut-off, very similar adjusted 5-  
17 and 10-year outcomes were observed within each risk group (1 vs 2-3 vs cN1) when  
18 using either STAMPEDE or conventional criteria (**Supplementary Table 6**,  
19 **Supplementary Figure 2**).

## 1    **DISCUSSION**

2    In this analysis comprising 3604 patients treated on 10 randomized trials of RT and  
3    ItADT for HRLPC, we noted statistically significant and clinically meaningful differences  
4    in long-term outcomes based on the overall number of baseline adverse risk factors.  
5    Specifically, patients with at least two risk factors (Gleason 8-10, cT3-4, PSA >20ng/mL)  
6    in context of cN0 disease, or cN1 disease (regardless of other risk factors) had poorer  
7    outcomes compared to those with only 1 risk factor, with a 5-year MFS of 78% for cN0  
8    patients with 2-3 risk factors and 67% for all patients with cN1 disease, versus 83% for  
9    patients with 1 risk factor and cN0. Moreover, the number of prostate cancer events  
10    contributing to the MFS and OS endpoints increased with the poorer risk groups,  
11    indicating that those patients more likely to develop life-threatening clinical events are  
12    potentially more likely to benefit from treatment intensification beyond RT and ItADT.

13    Since D'Amico and colleagues developed the first risk classification scheme for  
14    localized prostate cancer in the late 1990s,[12] the presence of biopsy Gleason 8-10,  
15    cT3-T4 and/or PSA >20ng/mL at diagnosis have been taken forward by guideline  
16    groups, such as EAU[4], ESMO[13] and NCCN[9], to define high-risk disease. However,  
17    outcomes within this group are heterogeneous and there have been subsequent efforts  
18    to refine risk stratification[14-17]. These have typically used these three variables to  
19    generate prognostic groups that are better able to risk-stratify patients, but have been  
20    limited by evaluation of patients undergoing surgery (and not RT and ADT),  
21    heterogeneity in treatments received and lack of significant numbers of patients  
22    receiving ItADT with RT. As such, our findings represent the largest study to define risk  
23    stratification within HRLPC, are the first to evaluate patients receiving ADT in addition to

1 RT, use IPD from randomized trials, and corroborate these earlier efforts that a simple  
2 assessment of the number of risk factors (1 vs 2-3 vs N1) can provide more robust  
3 prognostic information.

4 These results have several important implications for clinical practice as well as in the  
5 interpretation of ongoing (neo)adjuvant trials in HRLPC. The addition of 2 years of  
6 abiraterone to RT and ltADT has become a standard-of-care for “very” high-risk M0  
7 prostate cancer based on the STAMPEDE-abiraterone trial.[6] That comparison of the  
8 STAMPEDE study comprised of ~40% N1 patients (by conventional imaging), with the  
9 remainder having two of Gleason 8-10, cT3-4 or PSA  $\geq$ 40ng/mL, and the median PSA in  
10 the trial was 30-40ng/mL. In our analyses, very similar results in long-term outcomes  
11 were seen when patients were classified by either EAU/ESMO/NCCN high-risk criteria  
12 or STAMPEDE high-risk criteria. As such, N0 patients with 2 or 3 adverse risk factors  
13 (by EAU/ESMO/NCCN criteria) had a 5-year MFS <80% with RT and ltADT, and likely to  
14 benefit from the addition of abiraterone. In contrast, N0 patients with just one high-risk  
15 factor had better long-term outcomes with RT and ltADT, whereas N1 patients denoted  
16 a particularly high-risk group in whom intensification might be of greatest benefit.

17 There are several ongoing adjuvant trials assessing the addition of other ARPIs to RT  
18 and ltADT in HRLPC. Eligibility criteria vary between these trials, with baseline data  
19 from the ATLAS,[18] ENZARAD AND DASL trials[19] showing a range in cN1 disease  
20 from 11-28% and a median PSA in the ATLAS trial of 6ng/mL, which are notably  
21 different to the STAMPEDE population. Our results will be helpful to provide a  
22 framework upon which to guide clinical decision-making, based on extent of risk factors  
23 and by N0 vs N1 disease, thereby guiding the interpretation of these studies.



1 It is important to note that none of the patients included in our analysis had molecular  
2 imaging (e.g. PSMA-PET) for staging or evaluation of suspected recurrence or  
3 metastasis. PSMA-PET has greater sensitivity, specificity and diagnostic accuracy  
4 compared to conventional imaging in staging high-risk disease.[20] As such, our  
5 findings and outcome estimates only apply to those with high-risk and/or N1 disease on  
6 conventional scans, which is reflected in the 5-year MFS rate of 80% amongst high-risk  
7 N0 patients treated with RT and ItADT. This is lower than the 5-year MFS of 89% in  
8 patients with N0 disease treated with prostate-only RT and ItADT in the POP-RT trial,  
9 where the median PSA was similar to our cohort (28ng/mL vs. 24ng/mL), but 80% of  
10 patients were staged with PSMA-PET.[21] This indicates that the absence of nodal  
11 disease on PET is highly prognostic. As such, it is to be determined whether high-risk  
12 patients with one risk factor and <1cm PSMA-avid pelvic nodes (i.e. N0 by conventional  
13 imaging) would benefit from intensification of therapy beyond whole pelvis RT and ItADT  
14 alone.

15 The strengths of this work lie in the availability of IPD from multiple randomized trials  
16 with a median follow-up of nearly 9 years ensuring that the 5- and 10-year MFS  
17 estimates we provide are robust and can serve as a benchmark for ongoing trials and in  
18 counselling patients treated in routine practice. We specifically chose not to evaluate  
19 PSA-based endpoints, such as biochemical failure or event-free survival, since these  
20 have not shown to be good surrogates for OS.[22, 23] While there are other efforts  
21 ongoing to define which people may benefit most from addition of ADT (and beyond) to  
22 RT in high-risk disease,[24] the risk stratification we provide is based on inexpensive,  
23 readily available parameters that are already routinely used in everyday practice.

1 Despite these, we acknowledge key limitations, including the long time period over  
2 which trial participants were treated (1987-2016), lack of data on therapies utilized at  
3 recurrence, lack of data on the actual ADT duration that patients received, and  
4 heterogeneity in RT field, dose and fractionation, though we noted better outcomes  
5 amongst patients treated at RT doses of >70Gy (i.e. above the median) of this cohort, in  
6 line with recent data from the GETUG-AFU 18 study[25]. Nevertheless, we adjusted for  
7 RT dose and planned ADT duration as well as stratifying by years of enrolment in our  
8 multivariate analyses. We additionally lacked information on whether T staging was  
9 assigned by imaging or digital rectal exam (DRE), and outcomes might be better in  
10 those with radiologic T3-T4 disease only. Molecular imaging was not used in staging (or  
11 monitoring) patients, and studies are needed to define how PSMA-PET imaging can  
12 improve upon the data defined by clinicopathological variables and conventional  
13 imaging.

14 In summary, this IPD analysis comprising approximately 3600 patients treated with RT  
15 and ltADT for HRLPC demonstrated important prognostic differences between patients  
16 depending on the presence of specific risk factors (Gleason 8-10, cT3-4, PSA  
17 >20ng/mL; cN1), alone or in combination. Patients with 2-3 risk factors (in the context of  
18 cN0 disease) or cN1 disease (regardless of other risk factors) had 5-year MFS rates of  
19 <80% and appear to be the best candidates for intensification of therapy beyond RT and  
20 ltADT. These findings have implications for selection of patients for therapy  
21 intensification in clinical practice, and will be helpful in interpreting the results of ongoing  
22 adjuvant studies in HRLPC.

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## TABLES AND FIGURES

**Table 1 – Baseline characteristics at randomization of included patients**

	<b>N (%)</b>
Age, yrs, median (IQR)	68 (63-73)
Year of randomization	
1987-1994	724 (20)
1995-1999	256 (7.1)
2000-2004	768 (21)
2005-2009	850 (24)
2010-2016	1006 (28)
PSA at randomization, ng/mL, median (IQR)	24 (12-48)
<10	719 (20)
10-20	806 (22)
>20	2061 (57)
Unknown	18 (0.50)
Biopsy Gleason score	
<7	564 (16)
7	1069 (30)
8-10	1942 (54)
Unknown	29 (0.80)
Clinical T stage	
Tx1-2	1002 (28)

T3-4	2602 (72)
Clinical N1	422 (12)
Planned duration of ADT treatment	
18 months	365 (10)
≥24 months	3239 (90)
Radiotherapy dose, Gy, median (IQR)*	70 (69-74)

\* evaluable N=2990

Abbreviations: ADT-Androgen Deprivation Therapy; IQR – interquartile range

**Table 2 – Multivariable models estimating the associations between long-term outcomes and baseline clinical parameters**

	MFS		OS		TTM		PCSM	
	HR (95% CI)	p	HR (95% CI)	p	sHR (95% CI)	p	sHR (95% CI)	p
Biopsy Gleason $\geq 8$ (ref: $\leq 7$ )	1.45(1.29-1.63)	<.001	1.42(1.26-1.61)	<.001	1.84 (1.55-2.19)	<.001	2.08(1.66-2.60)	<.001
Clinical T3-4 (ref: Tx1-2)	1.28(1.13-1.45)	<.001	1.22(1.07-1.39)	0.003	1.58(1.30-1.91)	<.001	1.73(1.35-2.22)	<.001
PSA at randomization (ref: <10)								
10-20ng/mL	1.08(0.92-1.27)	0.4	1.05(0.89-1.25)	0.5	1.06(0.84-1.34)	0.6	1.03(0.77-1.38)	0.9
>20ng/mL	1.30(1.13-1.50)	<.001	1.21(1.05-1.41)	0.011	1.30(1.06-1.59)	0.011	1.05(0.82-1.36)	0.7
Clinical N1 (ref: N0)	1.86(1.56-2.21)	<.001	1.77(1.45-2.15)	<.001	2.17(1.73-2.73)	<.001	2.43(1.79-3.30)	<.001
Age at randomization (per year)	1.02(1.01-1.03)	<.001	1.04(1.03-1.05)	<.001	0.97(0.96-0.98)	<.001	0.97(0.96-0.99)	0.001
Radiotherapy dose (ref: $\leq 70$ Gy)								
>70 Gy	1.05(0.91-1.22)	0.5	1.00(0.86-1.17)	>0.9	0.96(0.78-1.18)	0.7	0.73(0.55-0.97)	0.032
Unknown	1.42(1.17-1.74)	0.001	1.35(1.09-1.68)	0.006	1.30(0.98-1.74)	0.071	1.19(0.82-1.74)	0.4
ADT $\geq 24$ months (ref: 18 months)	0.80(0.64-0.99)	0.039	0.93(0.74-1.18)	0.6	0.61(0.45-0.81)	0.001	0.73(0.49-1.08)	0.11

Abbreviations: ADT – androgen deprivation therapy; MFS – metastasis-free survival; OS – overall survival; TTM – time to metastasis; PCSM – prostate cancer-specific mortality; HR – hazard ratio; sHR – subdistribution hazard ratio; CI – confidence interval



**Table 3 – Unadjusted Kaplan Meier estimates of 5-year and 10-year MFS rates (95% CI) in various subgroups of patients, stratified by risk factors (Gleason score, PSA, cT stage; and cN1) at baseline. NB – all patients with cN1 disease were analyzed together and stratified by Gleason score at diagnosis.**

	Gleason 7		Gleason 8-10	
	Tx1-2	T3-4	Tx1-2	T3-4
5-year MFS				
PSA <10ng/mL	-	87 (82-91)	82 (76-87)	75 (69-80)
PSA 10-20ng/mL	-	81 (75-85)	84 (77-89)	79 (73-83)
PSA >20ng/mL	84 (79-87)	80 (76-83)	74 (67-79)	77 (73-80)
cN1	76 (67-82)		64 (58-69)	
10-year MFS				
PSA <10ng/mL		65 (57-72)	62 (54-68)	52 (43-60)
PSA 10-20ng/mL		57 (50-64)	63 (54-70)	59 (51-66)
PSA >20ng/mL	63 (57-68)	59 (54-64)	47 (39-54)	46 (40-52)
cN1	36 (20-53)		38 (28-47)	

Abbreviations: MFS – metastasis-free survival; CI – confidence interval

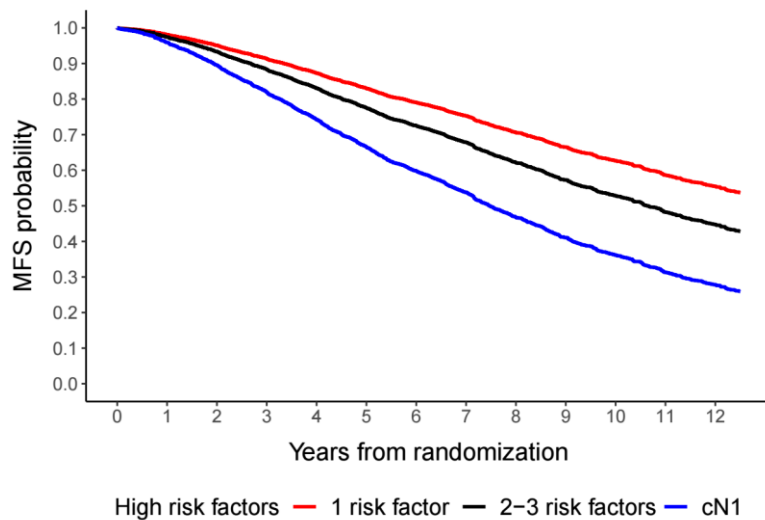
**Table 4 – Adjusted estimates of 5-year and 10-year MFS and OS from Cox regression and TTM and PCSM from the Fine and Gray models, based on number of baseline adverse risk factors (Gleason 8-10, cT3-4, PSA >20ng/mL) and cN1 disease.** All models were adjusted for age at randomization, ADT duration ( $\geq 24$  months vs 18 months) and radiotherapy dose ( $\leq 70$  Gy,  $>70$  Gy and unknown).

	<b>N</b>	<b>No. of events</b>	<b>5-year % (95% CI)</b>	<b>10-year % (95% CI)</b>
<b>MFS</b>				
1 risk factor	1241	508	83(81-85)	63(60-66)
2-3 risk factors	1900	796	78(76-79)	53(50-56)
cN1	422	188	67(62-71)	36(31-42)
<b>OS</b>				
1 risk factor	1241	467	87(86-88)	67(64-70)
2-3 risk factors	1900	683	84(82-85)	60(57-62)
cN1	422	144	77(74-80)	47(41-53)
<b>TTM</b>				
1 risk factor	1241	184	7.5(6.3-8.8)	15(13-17)
2-3 risk factors	1900	400	13(12-15)	25(23-28)
cN1	422	137	25(21-29)	44(38-50)
<b>PCSM</b>				
1 risk factor	1241	106	3.1(2.4-3.8)	8.0(6.6-9.6)
2-3 risk factors	1900	237	5.9(5.0-7.0)	15(13-17)
cN1	422	78	13(10-16)	30(25-35)

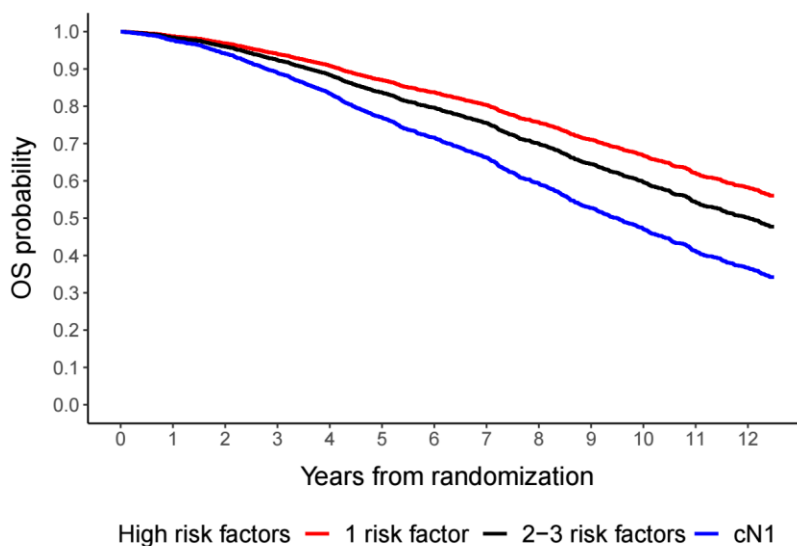
Abbreviations: MFS – metastasis-free survival; OS – overall survival; TTM – time to metastasis; PCSM – prostate cancer-specific mortality; CI – confidence interval

**Figure 1 – Adjusted curves showing MFS (2A) and OS (2B) from Cox regression models and TTM (2C) and PCSM (2D) from the Fine and Gray models, based on number of adverse baseline risk factors (Gleason  $\geq 8$ , cT3-4 and PSA  $>20\text{ng/mL}$ ) or cN1 disease. All models were adjusted for age at randomization, ADT duration ( $\geq 24$  months vs 18 months) and radiotherapy dose ( $\leq 70$  Gy,  $>70$  Gy and unknown).**

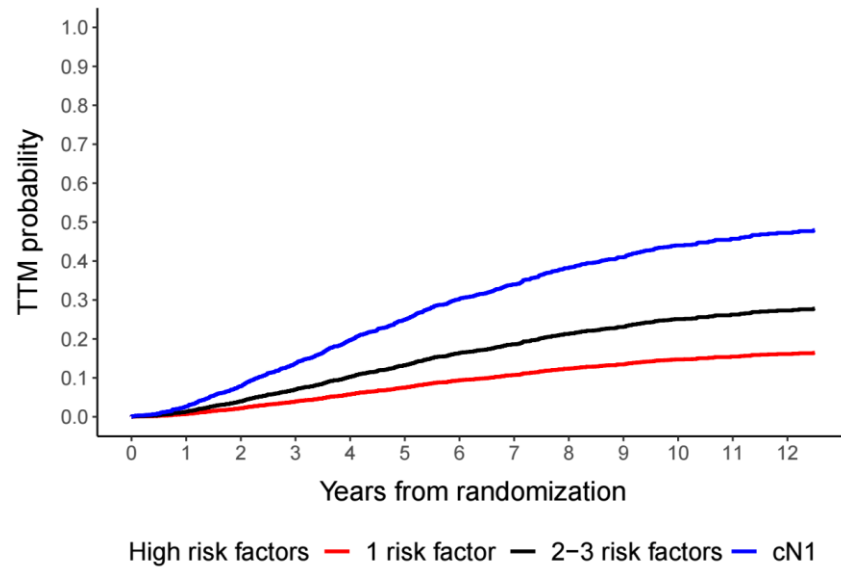
**1A:**



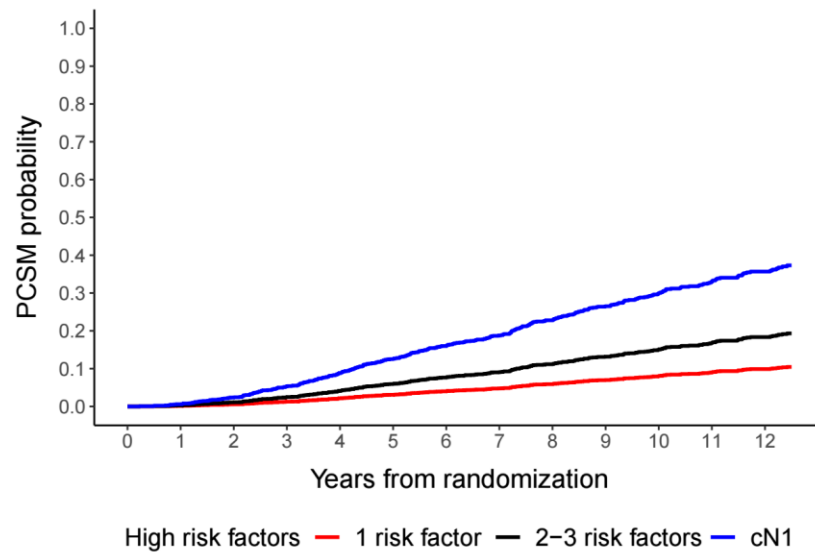
**1B**



**1C:**



**1D:**



Abbreviations: MFS – metastasis-free survival; TTM – time to metastasis; OS – overall survival; PCSM – prostate cancer-specific mortality

## SUPPLEMENTARY MATERIAL

**Supplementary Table 1 – Ongoing adjuvant trials in high-risk/locally advanced prostate cancer**

Trial (NCT)	N	Comparison	Eligibility	Results/Primary Endpoint
STAMPEDE (NCT00268476)	1974	Trial 1: ItADT vs ItADT + abiraterone  Trial 2: ItADT vs ItADT + abiraterone + enzalutamide	Either cN1 by conventional imaging or at least two of the following:  a) $\geq$ cT3  b) Gleason 8-10  c) PSA $\geq$ 40 ng/mL	MFS  Trial 1: HR=0.54 (0.43- 0.68)  Trial 2: HR=0.53 (0.39- 0.71)
ATLAS (NCT02531516)	1503	ItADT vs ItADT + apalutamide	1 of the following:  a) Gleason $\geq$ 8 and $\geq$ cT2c  b) Gleason $\geq$ 7, PSA $\geq$ 20 ng/mL, and $\geq$ cT2c	MFS
ENZARAD (NCT02446444)	802	ItADT vs ItADT + enzalutamide	1 of the following:  a) Gleason 8-10  b) Gleason 4+3 and $\geq$ cT2b-4 and PSA $\geq$ 20ng/mL  c) cN1 by conventional imaging	MFS
DASL-HiCAP (NCT04136353)	1100	ItADT vs ItADT + darolutamide	1 of the following:  a) Gleason 9-10  b) Gleason 8 and any of the following:  - $\geq$ cT2b	MFS

			<ul style="list-style-type: none"> <li>- MRI with T3a/T3b disease</li> <li>- PSA <math>\geq</math>20ng/mL</li> <li>c) cN1 by conventional imaging</li> </ul>	
PREDICT-RT (NCT04513717)	786	ltADT vs ltADT + apalutamide	Intensification arm: cN1 by conventional imaging, or Decipher >0.85 with 1 of the following: <ul style="list-style-type: none"> <li>a) PSA &gt;20ng/mL</li> <li>b) <math>\geq</math>cT3</li> <li>c) Gleason 8-10</li> </ul>	MFS
PEACE-2 (NCT01952223)	1048	Prostate +/- pelvic RT + ltADT +/- cabazitaxel	At least 2 of: <ul style="list-style-type: none"> <li>a) Gleason 8-10</li> <li>b) cT3-4</li> <li>c) PSA <math>\geq</math>20ng/mL</li> </ul>	cPFS

Abbreviations: ltADT – long-term ADT; RT – radiotherapy; cPFS – clinical progression-free survival

**Supplementary Table 2 – Trials included in the analysis**

Study	Year Enrolled	Arm	Treatment	RT dose, median (range)	Total N	Eligible N
EORTC 22863 <sup>1</sup>	1987-1995	Experimental	RT+ AADT 3yr	70 (18-90)	415	204
EORTC 22961 <sup>2</sup>	1997-2001	Experimental	RT + NADT 6mo + AADT 2.5yr	70 (64-74)	970	436
French study (Mottet) <sup>3</sup>	2000-2003	Experimental	RT + ADT 3yr	70 (65-76)	264	129
GICOR-DART 01/05 <sup>4</sup>	2005-2010	Experimental	RT + NADT 4mo + AADT 2yr	78 (31-83)	352	91
RTOG 9202 <sup>5</sup>	1992-1995	Experimental	RT + NADT 4mo + AADT 2yr	68 (13-77)	1520	617
GETUG 12 <sup>6</sup>	2002-2006	Control	RT + AADT 3yr	74 (69-80)	413	204
RTOG 0521 <sup>7</sup>	2005-2009	Control	RT + AADT 2yr	NA	563	281
RTOG 9902 <sup>8</sup>	2000-2004	Control	RT + AADT 2yr	NA	397	197
STAMPEDE <sup>9-12</sup>	2006-2016	Control	RT + AADT ≥2yr	74 (8.0-156)	2537	1080*
TROG 0304 <sup>13</sup>	2003-2007	Experimental	RT + NADT 6mo + AADT 12mo	70 (46-76)	1071	365
Total					8502	3604

\*In the STAMPEDE trial, there were 2142 controls from 1080 unique subjects for 7 treatment comparisons. If the same subject was used as multiple controls, data with longest follow-up was retained for individual patient level analysis.

Abbreviations: AADT – adjuvant ADT; NADT – neoadjuvant ADT; RT – radiotherapy; NA - Not available

References: <sup>1</sup> Bolla et al, Lancet 2002; <sup>2</sup> Bolla et al, NEJM 2009; <sup>3</sup> Mottet et al, Eur Urol 2012; <sup>4</sup> Zapatero et al, Lancet Oncol 2015; <sup>5</sup> Hanks et al, J Clin Oncol 2003; <sup>6</sup> Fizazi et al, Lancet 2015; <sup>7</sup> Rosenthal et al, J Clin Oncol 2019; <sup>8</sup> Rosenthal et al, Int J Radiat Oncol Biol Phys 2015; <sup>9</sup> James et al, Lancet 2016; <sup>10</sup> James et al, JNCI Cancer Spect 2022; <sup>11</sup> Mason et al, J Clin Oncol 2017; <sup>12</sup> Attard et al, Lancet 2022; <sup>13</sup> Denham et al, Lancet Oncol 2019

**Supplementary Table 3A – Unadjusted Kaplan Meier estimates of MFS and OS at 5- and 10-years in various risk subgroups**

Group	Total N	MFS			OS		
		No. of events	5-year % (95% CI)	10-year % (95% CI)	No. of events	5-year % (95% CI)	10-year % (95% CI)
GS≤7 T3-4 PSA<10	232	81	87 (82-91)	65 (57-72)	77	88 (82-91)	68 (60-75)
GS≤7 T3-4 PSA 10-20	286	127	81 (75-85)	57 (50-64)	120	84 (79-88)	64 (57-70)
GS≤7 Tx1-2 PSA>20	364	162	84 (79-87)	63 (57-68)	143	89 (85-92)	68 (62-73)
GS≤7 T3-4 PSA>20	612	253	80 (76-83)	59 (54-64)	221	84 (81-87)	66 (61-70)
GS ≤7 cN1	130	55	76 (67-82)	36 (20-53)	45	83 (75-88)	51 (33-66)
GS≥8 Tx1-2 PSA<10	195	74	82 (76-87)	62 (54-68)	70	89 (83-92)	64 (56-71)
GS≥8 Tx1-2 PSA 10-20	164	64	84 (77-89)	63 (54-70)	57	89 (83-93)	68 (59-75)
GS≥8 T3-4 PSA<10	231	95	75 (69-80)	52 (43-60)	79	82 (76-86)	59 (50-67)
GS≥8 T3-4 PSA 10-20	267	99	79 (73-83)	59 (51-66)	87	84 (79-88)	65 (57-72)
GS≥8 Tx1-2 PSA>20	220	116	74 (67-79)	47 (39-54)	104	84 (79-89)	53 (45-60)
GS≥8 T3-4 PSA>20	570	233	77 (73-80)	46 (40-52)	192	83 (79-86)	55 (49-61)
GS ≥8, cN1	289	132	64 (58-69)	38 (28-47)	98	76 (71-81)	46 (34-56)

Abbreviations: GS – Gleason score; MFS – metastasis-free survival; OS – overall survival



**Supplementary Table 3B – Unadjusted cumulative Incidence of TTM and PCSM at 5- and 10-years in various risk subgroups from competing risk models**

Group	Total N	TTM			PCSM		
		No. of events	5-year % (95% CI)	10-year % (95% CI)	No. of events	5-year % (95% CI)	10-year % (95% CI)
GS≤7 T3-4 PSA<10	232	20	4.1 (2.0-7.4)	9.3 (5.7-14)	15	3.3 (1.5-6.5)	7.0 (3.7-12)
GS≤7 T3-4 PSA 10-20	286	42	7.6 (4.8-11)	17 (12-22)	27	3.2 (1.6-5.8)	7.9 (4.8-12)
GS≤7 Tx1-2 PSA>20	364	60	7.1 (4.8-10)	16 (12-20)	23	1.4 (0.54-3.1)	6.3 (3.9-9.5)
GS≤7 T3-4 PSA>20	612	100	7.8 (5.8-10)	18 (14-22)	55	3.6 (2.3-5.3)	8.9 (6.3-12)
GS ≤7 cN1	130	38	19 (13-26)	46 (28-62)	25	11 (6.3-17)	31 (16-47)
GS≥8 Tx1-2 PSA<10	195	35	11 (6.8-16)	19 (13-25)	24	3.8 (1.7-7.3)	13 (8.4-19)
GS≥8 Tx1-2 PSA 10-20	164	27	8.9 (5.1-14)	18 (12-25)	17	3.3 (1.2-7.0)	11 (6.4-17)
GS≥8 T3-4 PSA<10	231	58	18 (13-23)	27 (21-34)	35	10 (6.8-15)	17 (11-23)
GS≥8 T3-4 PSA 10-20	267	56	14 (10-19)	24 (18-31)	41	7.9 (5.0-12)	18 (13-24)
GS≥8 Tx1-2 PSA>20	220	49	14 (10-19)	26 (20-32)	33	3.8 (1.8-7.0)	18 (12-24)
GS≥8 T3-4 PSA>20	570	137	15 (12-18)	33 (28-38)	73	6.7 (4.8-9.0)	19 (15-24)
GS ≥8, cN1	289	98	28 (23-33)	44 (35-53)	52	12 (8.8-16)	29 (21-38)

TTM – time to metastasis; PCSM – prostate cancer-specific mortality

**Supplementary Table 4 – Unadjusted estimates of 5-year and 10-year MFS, OS, TTM and PCSM based on number of baseline adverse risk factors (Gleason 8-10, cT3-4, PSA >20ng/mL) and cN1 disease, by age groups.**

	Age ≤68 years (median)				Age >68 years (median)			
	N	No. of events	5-year % (95% CI)	10-year % (95% CI)	N	No. of events	5-year % (95% CI)	10-year % (95% CI)
<b>MFS*</b>								
1 risk factor	611	215	85 (82-88)	67 (63-71)	629	293	81 (78-84)	56 (51-60)
2-3 risk factors	948	360	80 (77-83)	57 (52-61)	952	436	75 (72-78)	49 (45-54)
cN1	257	112	66 (60-72)	39 (27-51)	165	76	70 (62-76)	31 (18-45)
<b>OS*</b>								
1 risk factor	611	185	91 (88-93)	74 (69-77)	629	282	84 (81-87)	59 (54-63)
2-3 risk factors	948	288	87 (84-89)	66 (61-69)	952	395	80 (77-83)	55 (51-59)
cN1	257	80	79 (74-84)	52 (39-64)	165	64	77 (69-83)	37 (22-51)
<b>TTM**</b>								
1 risk factor	611	107	8.5 (6.4-11)	18 (15-22)	629	77	6.5 (4.7-8.7)	13 (10-16)
2-3 risk factors	948	236	14 (12-16)	30 (27-34)	952	164	12 (10-14)	20 (17-23)
cN1	257	92	29 (23-35)	49 (37-60)	165	45	19 (14-26)	41 (28-54)
<b>PCSM**</b>								
1 risk factor	611	59	2.6 (1.5-4.1)	9.2 (6.8-12)	629	47	3.0 (1.9-4.7)	8.0 (5.8-11)
2-3 risk factors	948	136	6.1 (4.7-7.8)	18 (15-21)	952	101	5.8 (4.4-7.5)	12 (9.8-15)
cN1	257	52	13 (9.0-17)	32 (21-44)	165	26	10 (6.3-16)	29 (17-42)

\* Kaplan-Meier estimates; \*\* Cumulative incidence from competing risk models

Abbreviations: MFS – metastasis-free survival; OS – overall survival; TTM – time to metastasis; PCSM – prostate cancer-specific mortality; CI – confidence interval

**Supplementary Table 5 – Unadjusted estimates of 5-year and 10-year MFS, OS, TTM and PCSM based on number of baseline adverse risk factors (Gleason 8-10, cT3-4, PSA >20ng/mL) and cN1 disease, by radiotherapy dose delivered.**

	Radiation dose ≤ 70 Gy				Radiation dose > 70 Gy			
	N	No. of events	5-year % (95% CI)	10-year % (95% CI)	N	No. of events	5-year % (95% CI)	10-year % (95% CI)
<b>MFS*</b>								
1 risk factor	659	297	83 (80-86)	59 (55-63)	282	109	83 (78-87)	63 (56-69)
2-3 risk factors	807	398	75 (72-78)	51 (47-55)	817	262	83 (80-85)	57 (51-62)
cN1	94	48	74 (64-82)	41 (26-55)	291	119	68 (62-73)	39 (27-51)
<b>OS*</b>								
1 risk factor	659	280	87 (84-90)	63 (58-67)	282	98	88 (83-91)	69 (62-74)
2-3 risk factors	807	363	80 (77-83)	57 (52-61)	817	204	88 (86-90)	66 (60-70)
cN1	94	44	83 (73-89)	49 (34-63)	291	84	79 (74-83)	49 (35-61)
<b>TTM**</b>								
1 risk factor	659	97	7.1 (5.2-9.2)	15 (12-19)	282	42	8.7 (5.7-13)	16 (12-22)
2-3 risk factors	807	180	13 (11-16)	25 (21-28)	817	143	11 (8.8-13)	23 (19-28)
cN1	94	31	21 (13-30)	41 (27-55)	291	89	25 (20-30)	44 (32-55)
<b>PCSM**</b>								
1 risk factor	659	58	2.5 (1.5-4.0)	9.5 (7.1-12)	282	25	3.1 (1.4-5.7)	8.1 (4.9-12)
2-3 risk factors	807	122	7.1 (5.4-9.0)	16 (13-19)	817	64	4.3 (3.0-5.8)	12 (8.6-15)
cN1	94	23	9.8 (4.8-17)	31 (19-44)	291	43	10 (7.2-14)	26 (17-37)

\* Kaplan-Meier estimates; \*\* Cumulative incidence from competing risk models

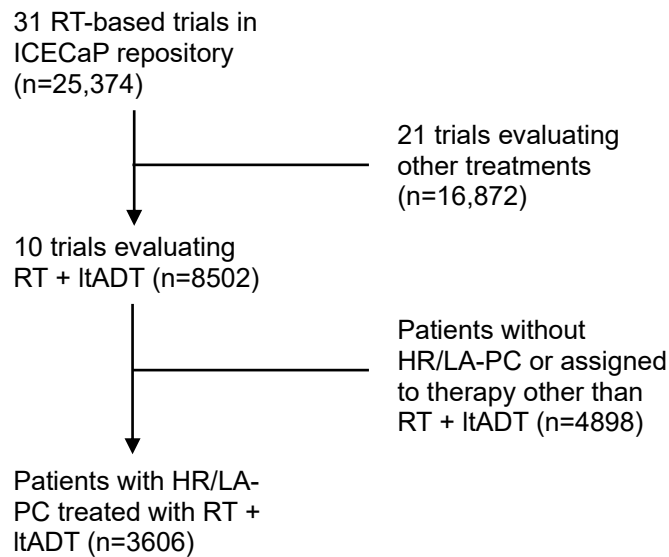
Abbreviations: MFS – metastasis-free survival; OS – overall survival; TTM – time to metastasis; PCSM – prostate cancer-specific mortality; CI – confidence interval

**Supplementary Table 6 – Adjusted estimates of 5-year and 10-year MFS and OS from Cox regression and TTM and PCSM from the Fine and Gray model, based on number of baseline adverse risk factors by the STAMPEDE criteria (Gleason 8-10, cT3-4, PSA  $\geq 40$ ng/mL) and cN1 disease.** All models were adjusted for age at randomization, ADT duration ( $\geq 24$  vs 18 months) and radiotherapy dose ( $\leq 70$  Gy,  $>70$  Gy and unknown).

	<b>N</b>	<b>No. of events</b>	<b>5-year % (95% CI)</b>	<b>10-year % (95% CI)</b>
<b>MFS</b>				
1 risk factor	1582	684	82(80-84)	61(58-63)
2-3 risk factors	1559	620	77(76-79)	53(50-56)
cN1	422	188	67(62-71)	36(31-42)
<b>OS</b>				
1 risk factor	1582	628	86(85-88)	65(62-68)
2-3 risk factors	1559	522	84(82-85)	60(57-63)
cN1	422	144	77(74-80)	47(41-53)
<b>TTM</b>				
1 risk factor	1582	241	7.7(6.6-8.7)	15(13-17)
2-3 risk factors	1559	343	14(13-16)	27(25-30)
cN1	422	137	25(21-29)	44(39-51)
<b>PCSM</b>				
1 risk factor	1582	142	3.2(2.6-3.9)	8.4(7.1-9.8)
2-3 risk factors	1559	201	6.5(5.5-7.6)	16(14-19)
cN1	422	78	13(10-16)	30(24-36)

Abbreviations: MFS – metastasis-free survival; OS – overall survival; TTM – time to metastasis; PCSM – prostate cancer-specific mortality; CI – confidence interval

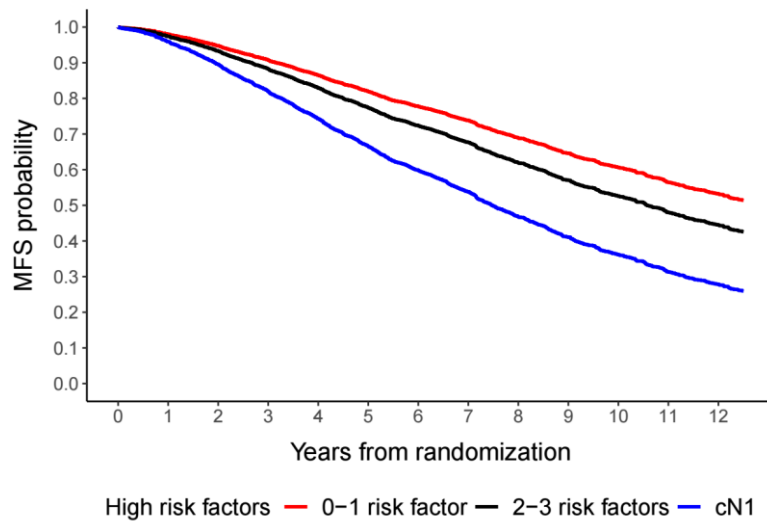
**Supplementary Figure 1 – Flowchart of selection of patients and trials included in the analysis**



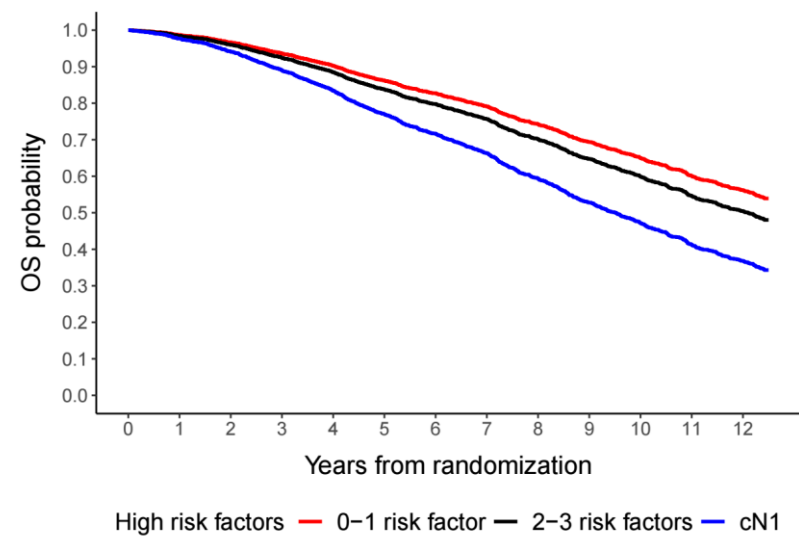
Abbreviations: RT – radiotherapy; ItADT – long-term ADT; HR/LA-PC – high-risk and/or locally-advanced prostate cancer

**Supplementary Figure 2 –Adjusted curves showing MFS (A) and OS (B) from Cox regression models and TTM (C) and PCSM (D) from the Fine and Gray models, based on number of adverse baseline risk factors by the STAMPEDE high-risk criteria (Gleason  $\geq 8$ , cT3-4 and PSA  $\geq 40$  ng/mL) or cN1 disease. All models were adjusted for age at randomization, ADT duration ( $\geq 24$  vs 18 months) and radiotherapy dose ( $\leq 70$  Gy,  $>70$  Gy and unknown).**

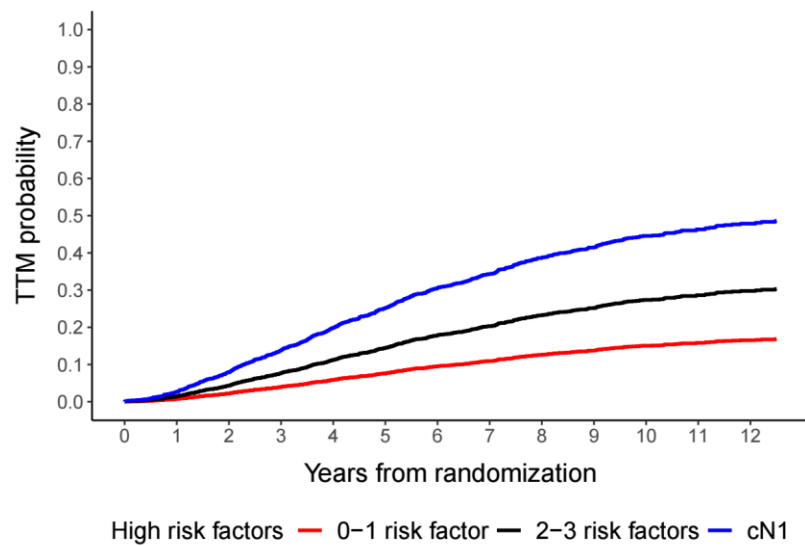
(A)



(B)



(C)



(D)

