GUIDELINES ON PROSTATE CANCER

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1 BACKGROUND

Cancer of the prostate (CaP) is now recognized as one of the principal medical problems facing the male population. In Europe, an estimated 2.6 million new cases of cancer are diagnosed each year. Prostate cancer constitutes about 11% of all male cancers in Europe (1), and accounts for 9% of all cancer deaths among men within the European Union (EU) (2). It is worth mentioning that there are comparatively large regional differences; for example, in Sweden, where there is a long life expectancy and a comparatively modest mortality from smoking-related diseases, CaP is the most common malignancy in males, accounting for 31.5% of all new cases of cancer in 1999 (3).

By the time of diagnosis, only 55% of tumours are clinically localized in the absence of an organized screening programme (4). Even in modern series, 30–45% of patients with clinically localized disease are found to have extracapsular extension at pathological staging (5,6).

1.1 REFERENCES


2 CLASSIFICATION

The 2002 TNM (Tumour Node Metastasis) classification for CaP is shown in Table 1 (1).

Table 1: Tumour Node Metastasis (TNM) classification of CaP

<table>
<thead>
<tr>
<th>T - Primary tumour</th>
<th>T2</th>
<th>Tumour confined within the prostate§</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>TX</td>
<td>Primary tumour cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>T0</td>
<td>No evidence of primary tumour</td>
</tr>
<tr>
<td>T1</td>
<td>T2</td>
<td>Tumour involves one half of one lobe or less</td>
</tr>
<tr>
<td>T1a</td>
<td>T2a</td>
<td>Tumour involves more than half of one lobe, but not both lobes</td>
</tr>
<tr>
<td>T1b</td>
<td>T2b</td>
<td>Tumour involves both lobes</td>
</tr>
<tr>
<td>T1c</td>
<td>T2c</td>
<td>Tumour identified by needle biopsy (e.g., because of elevated prostate-specific antigen (PSA) level)</td>
</tr>
<tr>
<td>T2</td>
<td>T3</td>
<td>Tumour extends through the prostatic capsule§</td>
</tr>
<tr>
<td>T2a</td>
<td>T3a</td>
<td>Extracapsular extension (unilateral or bilateral)</td>
</tr>
<tr>
<td>T2b</td>
<td>T3b</td>
<td>Tumour invades seminal vesicle(s)</td>
</tr>
<tr>
<td>T2c</td>
<td>T3c</td>
<td>Tumour identified by needle biopsy (e.g., because of elevated prostate-specific antigen (PSA) level)</td>
</tr>
<tr>
<td>T3</td>
<td>T4</td>
<td>Tumour is fixed or invades adjacent structures other than seminal vesicles: bladder neck, external sphincter, rectum, levator muscles, or pelvic wall</td>
</tr>
<tr>
<td>T3a</td>
<td>T4a</td>
<td>Tumour invades seminal vesicle(s)</td>
</tr>
<tr>
<td>T3b</td>
<td>T4b</td>
<td>Tumour invades adjacent structures other than seminal vesicles: bladder neck, external sphincter, rectum, levator muscles, or pelvic wall</td>
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</table>

N - Regional lymph nodes§

<table>
<thead>
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<th>N - Regional lymph nodes</th>
<th>N0</th>
<th>No regional lymph node metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>N1</td>
<td>N1</td>
<td>Regional lymph node metastasis</td>
</tr>
</tbody>
</table>

M - Distant metastasis§

<table>
<thead>
<tr>
<th>M - Distant metastasis</th>
<th>M0</th>
<th>No distant metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1</td>
<td>M1</td>
<td>Distant metastasis</td>
</tr>
<tr>
<td>M1a</td>
<td>M1a</td>
<td>Non-regional lymph node(s)</td>
</tr>
<tr>
<td>M1b</td>
<td>M1b</td>
<td>Bone(s)</td>
</tr>
<tr>
<td>M1c</td>
<td>M1c</td>
<td>Other site(s)</td>
</tr>
</tbody>
</table>

§ Tumour found in one or both lobes by needle biopsy, but not palpable or visible by imaging, is classified as T1c.

§ Invasion into the prostatic apex, or into (but not beyond) the prostate capsule, is not classified as T3, but as T2.

§ Metastasis no larger than 0.2 cm can be designated pN1mi.

§ When more than one site of metastasis is present, the most advanced category should be used.

2.1 Gleason score

The most commonly used system for grading adenocarcinoma of the prostate is the Gleason score (2). The system describes a score between 2 and 10, with 2 being the least aggressive and 10 the most aggressive. This score is the sum of the two most common patterns (grades 1-5) of tumour growth found. To be counted, a pattern (grade) needs to occupy more than 5% of the biopsy specimen. Biopsy material (core biopsy or operative specimens) is required to be able to assess the Gleason score; cytological preparations cannot be used.

Whenever possible, evidence levels and grade of recommendation have been inserted in this updated guidelines text according to the principles of evidence-based medicine (EBM) (3). (See Section 17 for full explanations).

2.2 REFERENCES


3 RISK FACTORS

The factors that determine the risk of developing clinical CaP are not well known, however, a few have been identified. An important risk factor seems to be heredity. If one first-line relative has the disease, the risk is at least doubled. If two or more first-line relatives are affected, the risk increases 5- to 11-fold (1,2). A small subpopulation of individuals with CaP (about 9%) has true hereditary CaP, defined as three or more relatives affected or at least two who have developed early-onset disease, i.e., before the age of 55 (3). The frequency of autopsy-detected cancers is roughly the same in different parts of the world (4). This finding is in sharp contrast with the incidence of clinical CaP, which differs widely between different geographical areas, being high in the USA and Northern Europe and low in South-East Asia (5,6). However, if Japanese men move from Japan to Hawaii, their risk of CaP increases, and if they move to California their risk increases even more and approaches that of American men (7).

These findings indicate that environmental factors affect the risk of progression from so-called latent CaP to clinical CaP. The identity of these factors is still under debate, but a high content of animal fat in the diet may be important in increasing the risk of developing CaP (8). Other factors include low intakes of vitamin E, selenium, lignans and isoflavonoids (9). It has also been proposed that the risk of developing clinical CaP is inversely related to sun exposure. Sunlight might be protective against CaP due to an increase in vitamin D levels (10).

In summary, hereditary factors are important in determining the risk of developing clinical CaP and environmental factors may have an important impact on this risk. The key question is whether or not there is enough evidence to recommend lifestyle changes (lowered intake of animal fat and increased intake of fruit, cereals and vegetables) in order to decrease the risk (11). There is some evidence for this, and this information could be given to male relatives of CaP patients who ask about the impact of diet (level of evidence: 3-4).

3.1 REFERENCES

4 SCREENING AND EARLY DETECTION

Population or mass screening is defined as the examination of asymptomatic men (at risk). Usually, screening takes place within the framework of a trial or study and is initiated by a screener. Contrary to that, early detection or opportunistic screening represents individual case findings. It is initiated by the screenee (patient) and/or his physician. The primary endpoint of both is two-fold: first, the reduction of CaP-specific mortality. The goal is not to detect more and more carcinomas nor is survival the endpoint because survival is heavily influenced by lead-time. Secondly, quality of life is important as expressed by quality of life adjusted gain in life years (QUALYs).

The trends in mortality from CaP show a wide variety from country to country all over the industrialized world (1). A decrease in mortality rates due to CaP is currently seen in the USA and Austria, but also in the UK and France, which share a similar decrease in CaP mortality rates (1). Similarly, in Sweden, the relative 5-year survival rates increased in the period from 1960 to 1988, which was attributed to increased diagnostic activities and the detection of more non-lethal tumours (2). However, this trend could not be confirmed in a similar study from the Netherlands (3).

The reduction in mortality seen lately in the USA is often attributed to the widely adopted aggressive screening policy. However, there is still no absolute proof that the concept of prostate-specific antigen (PSA) screening is the cause for reduced mortality due to CaP (4,5).

A non-randomized screening project in Tyrol (Austria) may support the hypothesis that screening can be effective in reducing CaP mortality. The early detection programme in combination with the availability of free treatment was used as an explanation for the 33% decrease in the CaP mortality rate seen in Tyrol as compared with the rest of Austria (6) (level of evidence: 2b). In addition, Labrie and co-workers from Quebec (Canada) claim lower mortality rates in men randomized to active CaP screening (7), even though these results have been challenged (8). Other studies have contradicted the positive findings attributed to screening, with a comparative study between the Seattle area (highly screened population) and Connecticut (seldom screened population) by Lu-Yao and coworkers (9) showing that, notwithstanding the very large diversity in PSA testing and in use of curative treatments, there was no difference in the reduction in the rate of CaP mortality (level of evidence: 2b).

In order to be able to really evaluate the efficacy of CaP screening, prospective, preferably population-based, randomized trials are needed. Two large trials are underway, the PLCO (Prostate, Lung, Colorectal and Ovary) trial in the USA and the ERSPC (European Randomized Screening for Prostate Cancer) in Europe (10). The first analysis of the main endpoint of these trials – differences in CaP mortality – is scheduled for 2008 (level of evidence: 1b).

Thus, at the present time, there is a lack of evidence to support or disregard widely adopted, population-based screening programmes for early detection of CaP aimed at all men in a given population (level of evidence: 3).

Less controversial, and recommended in most guidelines, is the use of PSA in combination with digital rectal examination (DRE) as an aid to early diagnosis (11) (see chapter 5) (level of evidence: 3).
4.1 REFERENCES


5 DIAGNOSIS

The main diagnostic tools used to look for evidence of CaP include DRE, serum concentration of PSA and transrectal ultrasonography (TRUS) (1). Diagnosis depends on the presence of adenocarcinoma in operative specimens, prostate biopsy cores or aspiration needle cytology. Histopathological examination also allows grading of the tumour. Multiple systematic ultrasound-guided biopsies will detect more cancers than digital- or ultrasound-guided biopsies of suspicious areas (2,3).

5.1 Digital rectal examination (DRE)
The majority of CaPs are located in the peripheral zone of the prostate and may be detected by DRE when the volume is about 0.2 mL or larger. The risk of a positive DRE turning out to be cancer is heavily dependent on the PSA value (Table 2) (4-6).

Table 2: PSA value and risk of CaP

<table>
<thead>
<tr>
<th>PSA ng/mL</th>
<th>PPV for cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-1</td>
<td>2.8-5%</td>
</tr>
<tr>
<td>1-2.5</td>
<td>10.5-14%</td>
</tr>
<tr>
<td>2.5-4</td>
<td>22-30%</td>
</tr>
<tr>
<td>4-10</td>
<td>41%</td>
</tr>
<tr>
<td>&gt; 10</td>
<td>69%</td>
</tr>
</tbody>
</table>

PPV = positive predictive value; PSA = prostate-specific antigen.

5.2 Prostate-specific antigen (PSA)
The measurement of PSA level has revolutionized the diagnosis of CaP (7). PSA is a kallikrein-like serine protease produced almost exclusively by the epithelial cells of the prostate. For practical purposes, it is organ-specific but not cancer-specific, and serum levels may be elevated in the presence of benign prostatic hypertrophy, prostatitis and other non-malignant conditions. PSA level, as an independent variable, is a better predictor of cancer than suspicious findings on DRE or TRUS (6).

Currently, many different commercial test kits for the measurement of PSA are available, but no common international standard exists (8). For the diagnosis of CaP, levels of other tumour markers, such as prostatic acid phosphatase (PAP) do not yield additional information if they are measured in addition to PSA (9). A threshold level of PSA that indicates the highest risk of CaP needs to be defined (10). The cumulative 7-year risk of being diagnosed with CaP in a screening programme based on PSA measurement only was 34% for men with PSA values between 3 and 6 ng/mL; 44% for those with PSA values between 6 and 10 ng/mL and 71% for those with PSA values > 10 ng/mL (11).

Thus, the detection of non-palpable CaP is dependent on the serum level of PSA. There is no universally accepted lower cut-off value, although > 4 ng/mL has been used in many studies. In younger men, aged 50-66 years, the CaP detection rate was 13.2% in the PSA interval 3-4 ng/mL; the majority of these cancers were judged to be clinically significant (12). Even lower cut-off levels have been proposed by some authors, still with a relatively high detection rate (13). The finding that many men may harbour CaP despite low levels of serum PSA has been underscored by the recent results from a US prevention study (14). The rate of CaP in relation to serum PSA for 2950 men in the placebo-arm and with normal PSA-values is presented in Table 3. The age range at biopsy was 62-91 years.

Table 3: Risk of CaP in relation to low PSA values

<table>
<thead>
<tr>
<th>PSA level (ng/mL)</th>
<th>Risk of CaP</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-0.5</td>
<td>6.6%</td>
</tr>
<tr>
<td>0.6-1</td>
<td>10.1%</td>
</tr>
<tr>
<td>1.1-2</td>
<td>17.0%</td>
</tr>
<tr>
<td>2.1-3</td>
<td>23.9%</td>
</tr>
<tr>
<td>3.1-4</td>
<td>26.9%</td>
</tr>
</tbody>
</table>

PSA = prostate-specific antigen.

These findings touch on an important issue associated with lowering the PSA level threshold and that is the avoidance of detecting insignificant cancers whose natural history is unlikely to be life-threatening (15).
Long-term data are not yet available from which to make a recommendation for the optimal PSA threshold value needed to detect non-palpable but clinically significant CaP (level of evidence: 3).

The following modifications of serum PSA value, which may improve the specificity of PSA in the early detection of CaP, have been described:

- PSA density (16)
- PSA density of the transition zone (17)
- Age-specific reference ranges (18)
- PSA molecular forms (19-21)
- PSA velocity (22)
- PSA doubling time (23).

All of the above modifications may help to distinguish between CaP and benign disorders of the prostate, particularly in the intermediate PSA range (4-10 ng/mL). Consensus has not been reached, however, on the application of these modifications in routine practice.

Stage T1c describes tumours recognized by biopsies performed because of an elevated PSA level only, with a normal DRE and TRUS. A review of the clinical relevance and pathological correlation of this tumour stage in European patients indicates that between 11% and 26% of cases are insignificant cancers, but between 18% and 49% represent locally advanced disease (24).

5.3 Transrectal ultrasonography (TRUS)
Different CaPs appear differently on TRUS. The classic picture of a hypoechoic area in the peripheral zone of the prostate will not always be seen (25). It must be stressed that many cancers are isoechoic and only detectable through systemic biopsies. Ellis and co-workers noted that 37.6% of their detected cancers were diagnosed in isoechoic areas of the prostate (3). Similar findings have been reported from several early detection studies.

TRUS has two potential roles in the diagnosis of CaP:
1. To identify lesions suspected of malignancy
2. To improve the accuracy of prostate biopsy.

It appears that, in a self-referred population, TRUS detects 50% more patients with CaP than physical examination (26,27). However, the ultrasonic appearance of CaP is variable, and it seems that only a very small number of cancers will be detected if DRE and PSA levels are normal (3,27,28). Thus, the main role of greyscale TRUS is to direct biopsies in order to obtain a systemic sampling of the gland.

5.4 Relationship between DRE, PSA, TRUS and CaP
The positive predictive value of various combinations of diagnostic procedures used in a screening population ranges from 20% to 80% (3,27,28). If a result using any one of the three modalities is abnormal, the positive biopsy rate is 6-25%; with two abnormalities it is 18-60%; and, if all three modalities are positive, it is 56-72%.

5.5 Prostate biopsies
Digitally guided fine-needle aspiration allows the diagnosis and cytological grading of the tumour with a minimal risk of complications (29). However, the method requires a specially trained cytologist to yield reproducible results and has never gained widespread use outside Scandinavian countries.

Ultrasound-guided transrectal 18G core biopsy has become the standard way to obtain material for histopathological examination. Multiple cores can be taken with a low risk of complications if antibiotic prophylaxis is used (30,31).

Lesion-guided biopsies can be used in cases where there is a palpable nodule in combination with a high PSA-level. Targeted biopsies directed by contrast-enhanced Doppler ultrasound have been shown to have a detection rate similar to that seen with systemic biopsies (32), but the method has not yet gained widespread acceptance (level of evidence: 2b).

Early studies indicated that the detection rate decreases when the number of cores decreases. Only patients with a PSA >10 ng/ml and a palpable nodule seemed to have an adequate detection rate with few, directed biopsies (33). For other patients, a more extensive sampling (more cores) were recommended.

Sextant biopsies, as described by Hodge et al., have been used in this situation (2). Lately, the standard way of obtaining sextant biopsies has been replaced by laterally directed sextant biopsies in order to optimize the CaP detection rate (34,35). Biopsy cores obtained this way include biopsies from the posterolateral aspect of the peripheral zone, the most common location for early CaP. The number of biopsies required for the optimal detection of CaP is controversial. Several studies have examined the detection rate with more biopsy cores at primary biopsy. Nearly all have shown a higher cancer detection rate in comparison
with the standard sextant technique described by Hodge. Eskew and co-workers, for instance, demonstrated that the five-region biopsy protocol with 13 to 18 cores increased the detection rate by 35% when compared to standard, mid-lobar sextant biopsies (36). Studies clearly show that the transition zone should not be the target area for a first set of prostate biopsies due to the consistently low cancer detection rate, which may be as low as 2% or less (37,38). The vast majority of the extra cancers were detected in the far-lateral mid-lobar region, an area well sampled by the technique of laterally directed sextant biopsy. It seems that the direction of the biopsies may well be as important as the number of cores.

If the first set of biopsies is negative, repeated biopsies can be recommended. In the second set of biopsies, a detection rate of about 10-35% has been reported in cases with a negative first set of biopsies (39-41). In cases where high-grade prostatic intraepithelial neoplasia (PIN) or atypical small acinar proliferation (ASAP) is present, as many as 50-100% of prostate cancers may be concomitant and re-biopsy is indicated (42-43). Djavan and co-workers found that two sets of biopsies detected the majority of clinically significant cancers (41). Even patients who have undergone more extensive biopsies may still have a significant detection rate at repeat biopsy (39,44). Today, we have no proven biopsy scheme that omits the need for re-biopsy in case of a persistent indication (level of evidence: 3).

With an increasing number of men undergoing more extensive biopsies at maybe two or even more occasions, the need for some form of analgesia has become more evident in clinical practice. Of the various methods examined, the use of a periprostatic injection with a local anaesthetic seems to combine high efficacy with easy application and low complication rates best. Of 23 studies, 20 have shown good efficacy when compared to placebo or intrarectal gel with local anaesthetic (45-67) (level of evidence:1a).

5.6 REFERENCES


34. Stamey TA. Making the most out of six systemic sextant biopsies. Urology 1995;45:2-12. 


38. Terris MK, Pham TQ, Issa MM, Kabalin JN. Routine transition zone and seminal vesicle biopsies in all patients undergoing transrectal ultrasound guided prostate biopsies are not indicated. J Urol 1997;157:204-206. 


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6. STAGING

The primary extension assessment of CaP is usually made by DRE, PSA measurement and bone scan, supplemented with computed tomography (CT)/magnetic resonance imaging (MRI) and chest X-ray in specific situations.

6.1 T-staging

The first level is the assessment of local tumour stage, where the distinction between intracapsular (T1-T2) and extracapsular (T3-T4) disease has the most profound impact on treatment decisions. DRE often underestimates the tumour extension, and in one study a positive correlation between DRE and pathological tumour stage was found in fewer than 50% of tumours (1). However, more extensive examinations for adequate T-staging are only recommended in selected cases when more precise staging directly affects the treatment decision, i.e., when curative treatment is an option.

Serum PSA levels increase with advancing stage. Nevertheless, when PSA level is measured in an individual patient it appears to have a limited ability to predict the final pathological stage accurately. Due to the production of PSA by benign and malignant prostatic tissue, there is no direct relationship between serum PSA concentration and the clinical and pathological tumour stage (2-4). A combination of serum PSA level, Gleason score on prostate biopsy and clinical T-stage, however, has proved to be more useful in predicting the final pathological stage than the individual parameters per se (5). The ability of the molecular forms of PSA to predict T-stage is still controversial. Although the free-to-total PSA ratio has been found to be useful for the staging of localized CaP (6), other studies have shown opposite results. Large multicentre studies are needed before any form of PSA can be used as a single modality for staging.

The most commonly used method for viewing the prostate is TRUS. However, only 60% of tumours are visible at TRUS and the remainder are not recognized due to their echogenicity. TRUS may reveal unsuspected extracapsular extension, but it does not determine tumour extent with sufficient accuracy to be
levels may indicate the presence of bony metastasis in 70% of affected patients (41). Furthermore, measurement of metastases accurately reflects the prognosis for an individual patient. Elevated skeletal alkaline phosphatase is a marker of the extent of skeletal metastases (8). The axial skeleton is involved in 85% of patients dying from CaP (40). The presence and extent of bone metastases is an important determinant of outcome in patients with CaP (51).  

6.3 M-staging

The axial skeleton is involved in 85% of patients dying from CaP (40). The presence and extent of bone metastases accurately reflect the prognosis for an individual patient. Elevated skeletal alkaline phosphatase levels may indicate the presence of bony metastasis in 70% of affected patients (41). Furthermore, measurement of metastases accurately reflects the prognosis for an individual patient. Elevated skeletal alkaline phosphatase is a marker of the extent of skeletal metastases (8). The axial skeleton is involved in 85% of patients dying from CaP (40). The presence and extent of bone metastases is an important determinant of outcome in patients with CaP (51).
of skeletal alkaline phosphatase and PSA at the same time increases clinical effectiveness to approximately 98% (42). In a prospective study, multiple regression analysis showed the extent of bone disease to be the only variable influencing the serum level of skeletal alkaline phosphatase and PSA.

However, in contrast to serum PSA, skeletal alkaline phosphatase demonstrated a statistical relationship with the extent of bone disease (43). Early detection of bone metastases will alert the clinician to the possible complications inherent in skeletal destruction. Bone scintigraphy remains the most sensitive method of assessing bone metastases, being superior to clinical evaluation, bone radiographs, serum alkaline phosphatase measurement and PAP determination (44,45). Technetium diphosphonates are the optimum radiopharmaceuticals currently available due to their extremely high bone-to-soft-tissue ratio (46). A semi-quantitative grading system based upon the extent of disease observed on the bone scan was found to correlate with survival (47). Besides bone, CaP may metastasize to any organ, but most commonly it affects distant lymph nodes, lung, liver, brain and skin. Clinical examination, chest X-ray, ultrasound, CT and MRI scans are all appropriate methods of investigation, but only if symptoms suggest the possibility of soft tissue metastasis.

The need for reliable serum markers to improve the pre-treatment staging of patients with CaP has long been recognized. At present, PSA is the marker of choice. A pre-treatment serum PSA level greater than 100 ng/mL was found to be the single most important indicator of metastatic disease, with a positive predictive value of 100% (48). Furthermore, it has helped to reduce the number of patients with newly diagnosed CaP who require a bone scan. Only on very rare occasions have patients with a low serum PSA concentration been found to harbour detectable skeletal metastases. The correlation between serum PSA and bone scintigraphy in patients with newly diagnosed untreated CaP has been further investigated (49-53). Results suggest that a staging bone scan may be superfluous if the serum PSA concentration is less than 20 ng/mL in asymptomatic patients with well- or moderately differentiated tumours. In contrast, patients with poorly differentiated tumours and locally advanced disease a staging bone scan should be obtained irrespective of the serum PSA value (54,55).

6.4 GUIDELINES FOR DIAGNOSIS AND STAGING OF CAP

1. An abnormal DRE result or elevated serum PSA measurement may indicate CaP. The exact cut-off level of what is considered to be a normal PSA value has not yet been determined, but values around < 2 – 3 ng/mL are often used for younger men (grade C recommendation)

2. The diagnosis of CaP depends on histopathological (or cytological) confirmation (grade B recommendation). Biopsy and further staging investigations are only indicated if they affect the management of the patient (grade C recommendation)

3. Transrectal ultrasound guided systemic biopsies is the recommended method in most cases with the suspicion of CaP. A minimum of 6-10 systemic, laterally directed, cores are recommended, eventually with more cores in larger glands (grade B recommendation):
   • Transition zone biopsies are not recommended in the first set of biopsies due to low detection rates (grade C recommendation)
   • One set of repeat biopsies are warranted in cases with persistent indication (abnormal DRE, elevated PSA or histopathological findings suggestive of malignancy at the initial biopsy) for prostate biopsy (grade B recommendation)
   • Overall recommendations for further (third or more) sets of biopsies cannot be made; the decision has to be made based on an individual patient (grade C recommendation)

4. Transrectal periprostatic injection with a local anaesthetic may be offered to patients as effective analgesia when undergoing prostate biopsies (grade A recommendation)

5. Local staging (T-staging) of CaP is based on findings from DRE and possibly MRI. Further information is provided by the number and sites of positive prostate biopsies, tumour grade and level of serum PSA (grade C recommendation)

6. Lymph node status (N-staging) is only important when potentially curative treatment is planned for. Patients with stage T2 or less, PSA < 20 ng/mL and a Gleason score ≤ 6 have less than a 10% likelihood of having node metastases and may be spared nodal evaluation. Accurate lymph node staging can only be determined by operative lymphadenectomy (grade B recommendation)

7. Skeletal metastasis (M-staging) is best assessed by bone scan. This may not be indicated in asymptomatic patients if the serum PSA level is less than 20 ng/mL in the presence of well- or moderately differentiated tumours (grade B recommendation).
6.5 REFERENCES


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7 TREATMENT: DEFERRED TREATMENT
(WATCHFUL WAITING; ACTIVE MONITORING)

7.1 Introduction
7.1.1 Definition
The term deferred treatment or watchful waiting (WW) is used to describe a treatment strategy that includes an active standpoint to postpone treatment until it is required. This does not only mean that treatments, such as palliative or hormonal, are withdrawn until symptomatic progression occurs (local or systemic). In rare, selected cases, this approach may also include younger patients with localized disease for whom potentially curative treatments are withheld until an indication for tumour activity occurs (i.e., rising serum PSA level, deteriorating histopathological factors on repeat biopsy). Patients who are offered WW must be followed-up carefully. It is worth mentioning that a patient’s anxiety is also a symptom that might warrant active treatment.

7.2 Deferred treatment of localized CaP (stage T1-T2, Nx-N0, M0)
There have been several attempts to summarize the key papers dealing with deferred treatment in patients with presumed localized CaP (1-6). Most of these papers present the same results as they analyze roughly the same series, but with a somewhat different methodology.

The paper by Chodak and co-workers is a pooled analysis of the original data from 828 patients treated by WW (1). It is based on patients from six non-randomized studies (6-13). The results describe cancer-specific survival and metastasis-free survival after 5 and 10 years of follow-up (1) (level of evidence: 2b).

The importance of tumour grade is clear, with very low survival rates for grade 3 tumours. Even if the 10-year cancer-specific survival rate is equally good (87%) for grade 1 and 2 tumours, the latter have a significantly higher progression rate, with 42% of the patients having developed metastases (Table 4).

Table 4: Outcome of deferred treatment in localized CaP in relation to tumour grade (1)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Disease-specific survival</th>
<th>Percentage of patients (95% confidence interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>5 years</td>
</tr>
<tr>
<td>Grade 1</td>
<td>98 (96-99)</td>
<td>87 (81-91)</td>
</tr>
<tr>
<td>Grade 2</td>
<td>97 (93-98)</td>
<td>87 (80-92)</td>
</tr>
<tr>
<td>Grade 3</td>
<td>67 (51-79)</td>
<td>34 (19-50)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Grade</th>
<th>Metastasis-free survival</th>
<th>Percentage of patients (95% confidence interval)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>5 years</td>
</tr>
<tr>
<td>Grade 1</td>
<td>93 (90-95)</td>
<td>81 (75-86)</td>
</tr>
<tr>
<td>Grade 2</td>
<td>84 (79-89)</td>
<td>58 (49-66)</td>
</tr>
<tr>
<td>Grade 3</td>
<td>51 (36-64)</td>
<td>28 (13-41)</td>
</tr>
</tbody>
</table>

The importance of tumour grade on survival after conservative management of CaP was also underlined in a large register study utilizing the Surveillance, Epidemiology, and End Results (SEER) database of the National Cancer Institute in the USA (14) (level of evidence: 3). Patients with grade 1, 2 and 3 tumours had 10-year cancer-specific survival rates of 92%, 76% and 43%, respectively, in agreement with the data from the pooled analysis.
The paper by Chodak and co-workers also specifically described the outcome for stage T1a patients (1), with cancer-specific 10-year survival rates of 96% and 94%, respectively, for grade 1 and 2 tumours. The metastasis-free survival rate was 92% for patients with grade 1 tumours, but 78% for those with grade 2 tumours, indicating a higher risk of progression in individuals with moderately differentiated tumours. This difference in progression rate is in accordance with other studies on stage T1a disease (15,16). To stage patients accurately and not overlook the presence of more extensive and/or more poorly differentiated tumours, repeat examinations with PSA measurement, TRUS and needle biopsy of the prostatic remnant have been advocated, especially in younger males with a long life expectancy (17).

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The impact of grade on the risk of tumour progression and ultimately death from CaP is also described in a paper by Albertsen and co-workers (18). They re-evaluated all biopsy specimens using the more widely accepted Gleason score and showed that the risk of CaP death was very high in Gleason 7-10 tumours, intermediate in Gleason 6 tumours, but low in Gleason 2-5 cancers (Table 5) (18,19) (level of evidence: 3). This paper also shows that Gleason 6-10 tumours carry a continuously increasing risk of ending the patient's life for up to 15 years of follow-up after conservative management. The cancer-specific survival curves for this group of patients have been published in a recent discussion article on different methods to assess outcome in treatment for localized CaP (19).

Table 5: The 15-year risk of dying from CaP in relation to Gleason score at diagnosis in patients with localized disease aged 55-74 years (17,18)

<table>
<thead>
<tr>
<th>Gleason score</th>
<th>Risk of cancer death</th>
<th>Cancer-specific mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-4</td>
<td>4-7%</td>
<td>8%</td>
</tr>
<tr>
<td>5</td>
<td>6-11%</td>
<td>14%</td>
</tr>
<tr>
<td>6</td>
<td>18-30%</td>
<td>44%</td>
</tr>
<tr>
<td>7</td>
<td>42-70%</td>
<td>76%</td>
</tr>
<tr>
<td>8-10</td>
<td>60-87%</td>
<td>93%</td>
</tr>
</tbody>
</table>

1 The figures on the risk of cancer death differ for different age groups and represent the true risk (taking actual competing mortality from other causes into consideration) in the studied population. The cancer-specific mortality compensates for differences in competing mortality and indicates the outcome if the patient actually lived for 15 years.

Data that include PSA and PSA changes over time are relatively sparse in the literature. In a recent review article, it was pointed out that patients with a PSA of < 3 ng/mL had no mortality from CaP within the first 10 years and that PSA changes over time were relatively unreliable in determining the risk for tumour progression (20).

The data above indicate a high risk of tumour progression after conservative treatment for some patients with apparently localized CaP. This has been supported by the results of other studies in which patients with a life expectancy exceeding 10 years have been shown to have a higher mortality rate from CaP when left without curative treatment (21-23). Long-term follow-up of the Johansson series renders the same outcome; a higher risk of dying from CaP in patients surviving more than 15 years with well and moderately differentiated tumours at diagnosis (24) (level of evidence: 3). Recently, a prospective randomized controlled trial comparing radical prostatectomy with conservative management showed a significant reduction in disease-specific mortality for the treatment group assigned to surgery (25) (level of evidence: 1b). For patients who choose deferred treatment, the risk of delaying hormonal therapy until disease progression occurs seems modest, although shorter cancer-specific survival times have been reported after deferred therapy compared with immediate hormonal therapy in presumed localized CaP (not utilising PSA for staging) after 15 years of follow-up (26). In contradiction to the finding from Lundgren, et al., a recent report from the EPC programme showed that a higher mortality was noted in the group of men with localized CaP treated with bicalutamide 150 mg compared to those who received placebo (27).

In summary, it seems that hormonal therapy should be withheld until there is definitive proof of disease activity (progression) but one may speculate whether there is some benefit to deliver it before the patient develops metastatic disease (see below).

### 7.3 Deferred treatment for locally advanced CaP (stage T3-T4, Nx-N0, M0)

The literature reporting on deferred treatment for locally advanced CaP is sparse. There are no randomized studies that compare more aggressive treatments, such as radiation therapy or surgery, eventually in combination with hormones. Most patients whose disease progresses after deferred treatment of locally advanced CaP will be candidates for hormonal therapy. There are reports from non-randomized studies showing that hormonal treatment may safely be delayed until metastatic progression occurs, as no survival advantage was noted between patients treated with immediate orchiectomy compared with delayed treatment.
However, when early and delayed treatments were compared in a large randomized trial carried out by the Medical Research Council (MRC), a survival benefit for immediate hormonal therapy was demonstrated (30), comparable with the results of Lundgren, et al. mentioned earlier (26) (level of evidence: 1b). Also, comparing placebo with bicalutamide 150 mg showed that in patients with locally advanced CaP, progression free survival was better with early treatment (27) (level of evidence: 1b).

Fifty selected asymptomatic patients (mean age 71 years) with highly or moderately differentiated stage T3 M0 CaP were followed up for 169 months (31). The 5- and 10-year cancer-specific survival rates were 90% and 74%, respectively, and the likelihood of being without treatment at 5 and 10 years was 40% and 30%, respectively. The authors concluded that WW may be a treatment option for selected patients with non-poorly differentiated T3 tumours and a life expectancy of less than 10 years (level of evidence: 3).

7.4 Deferred treatment for metastatic CaP (stage M1)
There are only very sparse data on this subject. The only candidates for such treatment should be asymptomatic patients with a strong wish to avoid treatment-related side-effects (level of evidence: 4). As the median survival time is about 2 years, the time without treatment (before symptoms occur) is very short in most cases. The MRC trial highlighted the risk of developing symptoms (pathological fractures, spinal cord compression) and even death from CaP, without receiving the possible benefit from hormonal treatment (30,32) (level of evidence:1b). If a deferred treatment policy is chosen for the patient with advanced CaP, there must be a possibility of close follow-up.

7.5 SUMMARY ON DEFERRED TREATMENT
7.5.1 Indications
In presumed localized CaP (Nx-N0, M0):
• Stage T1a - well and moderately differentiated tumours. In younger patients with a life expectancy of > 10 years, re-evaluation with PSA, TRUS and biopsies of the prostatic remnant is recommended (level of evidence: 2a)
• Stage T1b-T2b - well and moderately differentiated tumours. In asymptomatic patients with a life expectancy of < 10 years (level of evidence: 2a).

7.5.2 Options
In presumed localized CaP (Nx-N0, M0):
• Stage T1b-T2b patients, who are well informed and have well-differentiated, or Gleason 2-4, CaP and a life expectancy of 10-15 years
• All patients not willing to accept side-effects of active treatment
• Well-informed, asymptomatic patients with high PSA levels for whom cure is unlikely (level of evidence: 3)

In locally advanced disease (stage T3-T4):
• Asymptomatic patients with well or moderately differentiated cancer, CaP and a short life expectancy (level of evidence: 3)

In metastatic disease (M1):
• A very rare patient without any symptoms and the possibility of close follow-up (level of evidence: 4).

7.6 REFERENCES


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8 TREATMENT: RADICAL PROSTATECTOMY

8.1 Introduction
The surgical treatment of CaP consists of radical prostatectomy, meaning the removal of the entire prostate gland between urethra and bladder, with resection of both seminal vesicles. The procedure is routinely performed either retroperitoneally or using a transperineal approach. A few centres have gained experience with laparoscopic radical prostatectomy (1,2,3).

Radical prostatovesiculectomy was first applied at the beginning of the 20th century by Young (4) who used a perineal approach, while Memmelaar and Millin performed retropubic radical prostatectomy for the first time (5). In 1982, Walsh and Donker described the anatomy of the dorsal venous complex and the neurovascular bundles. This resulted in a significant reduction of the blood loss and improved continence and potency rates (6).

Currently, radical prostatectomy is the only treatment for localized CaP that has shown a cancer-specific survival benefit when compared to conservative management in a prospective, randomized trial (7). Surgical expertise has decreased the complication rates and improved cancer cure (8). In the hands of an experienced urological surgeon, the procedure is associated with minimal intra-operative and postoperative morbidity (9-11).

The retropubic approach is more commonly performed, as it enables simultaneous pelvic lymph node assessment to be carried out – an advantage over the perineal approach. It has been suggested that perineal radical prostatectomy might result in positive surgical margins more often than the retropubic approach (12), but this has not been confirmed (13). It is likely that laparoscopic lymphadenectomy and perineal prostatectomy have lower morbidity than the retropubic operation, but randomized studies are as yet unavailable. In recent years, some European centres have gained considerable experience with laparoscopic prostatectomy, and although long-term data on oncological outcome and complications are still lacking, this approach is becoming more and more accepted (14).

In men with localized CaP and a life expectancy of 10 years or more, the goal of a radical prostatectomy by any approach must be eradication of the disease (15). In fact, there is no rigid age limit for radical prostatectomy and a patient should not be denied this procedure on the grounds of age alone (16). However, it is worth pointing out that increasing co-morbidity with increasing age substantially decreases the actual risk of dying from localized CaP in men over the age of 70 years (17).

8.2 Stage T1a-T1b CaP
Stage T1a CaP is an incidental histological finding of cancer in 5% or less of resected prostatic tissue (transurethral resection of the prostate [TURP] or open adenomectomy), while it is a T1b stage when more than 5% contains cancer, or when the tumour is poorly differentiated. Although the risk of disease progression of untreated T1a CaP after 5 years is only 5%, these cancers can progress in about 50% of cases after 10-13 years (18). Thus, in younger patients with a life-expectancy of 15 years or more, the chance of disease progression is real, especially when a higher grade tumour is present.

In contrast, most patients with T1b tumours are expected to show disease progression after 5 years and aggressive treatment is often warranted (18). Consequently, it is very important to distinguish between T1a and T1b tumours and systematic prostate puncture biopsy of the remnant prostate 3 months after surgery is useful. Patients with T1b lesions are offered radical prostatectomy when they have a life expectancy of 10 years or more. Radical prostatectomy can become very difficult after a thorough TURP when almost no residual prostate is left behind (19), and external beam radiotherapy can be a valuable alternative treatment modality.

8.3 Stage T1c CaP
The clinically inapparent tumour identified by needle biopsy because of an aberrant PSA level has become the most frequent clinical stage in the actual radical prostatectomy population. In an individual patient, it is difficult to differentiate between clinically insignificant and life-threatening CaP. Most reports, however, stress that
PSA-detected tumours are mostly significant and should not be left untreated since up to 30% of T1c tumours are locally advanced (20). The proportion of insignificant tumours detected because of PSA elevation varies between 11% and 16% (21,22). Increasing the number of biopsies (sexant, octant...) might carry the risk of detecting more insignificant cancers but as shown in a recent study, increasing the number of biopsies to 12 did not increase the number of detected insignificant tumours (23).

The occurrence of PIN is not considered to be an indication for treatment, although 30% of patients with high-grade PIN will present an invasive adenocarcinoma within 5 years and 80% within 10 years (24). However, recently, low-grade PIN was shown not to be that inoffensive and the detection rate of invasive carcinoma in subsequent sets of biopsies was overall comparable to that in patients with high-grade PIN (25). Nevertheless, without proof of an invasive carcinoma, radical prostatectomy is not indicated because PIN may be a reversible phenomenon (26).

The major problem is how to recognize tumours on prostate puncture biopsy that do not need radical prostatectomy as they will be insignificant on the definitive pathological examination of the resected specimen. The needle biopsy findings and the free PSA ratio help in predicting insignificant disease (27). Using the Partin tables (which were updated in 2001) may be very helpful to determine the final pathological stage in order to better select patients that need to be surgically treated (28). Other authors have suggested the incorporation of biopsy information such as the number of cores or the percentage of cores invaded (29). When only one or a few cores are invaded and when the percentage of invasion in one core is limited, the chance of finding an insignificant CaP is more likely, certainly when the lesion is of low Gleason grade (30). It might be reasonable to follow-up some patients whose tumours are most likely to be insignificant. In general, however, radical prostatectomy should be advocated for patients with T1c tumours, keeping in mind that significant tumours will be found in the majority of these individuals.

8.4 Stage T2 CaP

Radical prostatectomy is one of the recommended standard treatments for patients with stage T2 CaP and a life expectancy of more than 10 years (31). The prognosis is excellent when the tumour is confined to the prostate based on pathological examination (32,33). Although most poorly differentiated tumours extend outside the prostate, patients with high-grade tumours that are confined to the prostate at histopathological examination still have a good prognosis after radical prostatectomy (34). A watchful waiting policy has been proposed for T2 tumours (35). If WW is proposed for low-grade T2 cancer, it should be remembered that pre-operative assessment of tumour grade by needle biopsy is frequently unreliable (36).

When the tumour is palpable or visible and clinically still confined to the prostate, disease progression can be expected in most patients who are long-term survivors. The median time to progression of untreated T2 disease is reported to be 6-10 years. T2a patients with a 10-year life expectancy should be offered radical prostatectomy since 35-55% of them will have disease progression after 5 years if not treated. T2b cancer still confined to the prostate but involving more than half of a lobe or both lobes will progress in more than 70% of patients within 5 years (37). These data have been confirmed by a large randomized trial comparing radical prostatectomy and watchful waiting that included mostly T2 CaP patients showing a significant reduction in disease-specific mortality (7).

In young men with localized CaP who are otherwise healthy, total surgical removal is an excellent option, and if it is performed by an experienced surgeon, the patient's subsequent quality of life should be more satisfactory. However, in an older patient or one with clinically significant co-existing conditions, radiation therapy could be the best option (38).

Lower rates of positive surgical margins for high-volume surgeons suggest that experienced and careful attention to surgical details, adjusted for the characteristics of a cancer being treated can decrease positive surgical margin rates and improve cancer control with radical prostatectomy (39).

8.5 Stage T3 CaP

T3a cancer is defined as capsular perforation and T3b cancer as invasion of the seminal vesicles. In the past, locally advanced CaP was seen in about 40% of all clinically diagnosed tumours. This figure must be lower today, but its management remains as controversial. In extracapsular tumours, radical prostatectomy often results in incomplete tumour excision. Higher morbidity and a substantially higher risk of local disease recurrence may be associated with those tumours compared with those confined to the prostate. In most patients, disease will finally progress systemically. Whether or not T3 CaP should be considered an indication for surgical treatment has therefore been questioned. The published reports on treatment outcomes in patients with clinical T3 are few (40-47).

Surgical treatment of clinical stage T3 CaP is traditionally discouraged (48), mainly because patients have an increased risk of positive surgical margins and both lymph node metastases or distant relapse (49,50). Combination treatment with hormonal and radiation therapy is gaining popularity, although it has not been demonstrated that this approach is superior to surgical treatment. A randomized study on radiotherapy with
hormones versus radiotherapy alone showed a clear advantage for the combination treatment, but did not show the superiority over radical prostatectomy (51). Another problem is “contamination” by the additional use of either adjuvant radiotherapy or immediate or delayed hormonal treatment in most of the series that reported on the treatment of clinical T3 CaP.

In the absence of data from randomized clinical trials comparing possible options for definitive therapy in these patients, only single or multicentre reports can be used to define the role of radical prostatectomy in this stage. Most studies have demonstrated that about 15% of all clinical stage T3 tumours were overstaged (cT3, pT2), while only 8% were understaged (cT3, pT4) (41). Patients who were overstaged obviously did very well, while most of those with pT3b cancer showed early disease progression.

For clinical T3 cancer, the overall PSA-free survival rate is about 20% after 5 years. The Gleason score of the tumour has a definite impact on progression (34), but there is not always a reliable correlation between the biopsy and the specimen Gleason score. On the other hand, seminal vesicle invasion, lymph node invasion, positive surgical margins and high PSA level are independent prognostic factors of PSA-free survival. Some authors have used a serum PSA level of 25 ng/mL as the discriminator for outcome (27,45). Others have shown that radical prostatectomy for clinical T3a cancer with a PSA below 10 ng/mL can achieve a 5-year PSA-free survival rate exceeding 60% (46).

Therefore, surgery has to be considered a therapeutic option for some patients with clinical T3a CaP. Not only clinically overstaged patients (pT2), but also individuals whose tumours actually are pT3a, can benefit from this treatment option. The problem remains in selecting those patients before surgery who have no lymph node involvement or seminal vesicle invasion. Nomograms, including PSA level, stage and Gleason score, can be useful in predicting the pathological stage of disease (28). In addition, nodal imaging with CT scan and seminal vesicle imaging with MRI or directed specific puncture biopsies of the nodes or of the seminal vesicles can be helpful in recognizing those patients who would not benefit from a surgical approach (52). Radical prostatectomy for clinical T3 cancer necessitates sufficient surgical expertise in order to keep the level of morbidity acceptable. Increased overall surgical experience must contribute to a decreased operative morbidity and to better functional results after radical prostatectomy for clinical T3 cancer.

### 8.6 Nodal disease

The indication for radical prostatectomy in all previously described stages assumes the pathologically proven absence of nodal involvement. Lymph node-positive (N+) disease will mostly be followed by systemic disease progression and all patients with significant N+ disease will ultimately fail. Nevertheless, the combination of radical prostatectomy and simultaneous hormonal treatment has been shown to achieve a 10-year cancer-specific survival rate of 80% (53). However, it is questionable whether or not these results could be obtained with hormonal treatment alone.

Most urologists are reluctant to perform radical prostatectomy for clinical N+ disease or will cancel surgery if a frozen section shows lymph node invasion. It should be noted that the definitive pathological examination after radical prostatectomy can show microscopic lymph node invasion. The incidence of tumour progression is lower in patients with fewer positive lymph nodes and in those with microscopic invasion only. N+ patients usually have significant nodal involvement and will be treated with hormonal manipulation only. In patients who prove to be pN+ after radical prostatectomy, adjuvant hormonal treatment can be advocated, but the benefits should be judged against side-effects of long-term hormonal therapy. PSA follow-up and hormonal treatment in case of PSA rise is therefore an acceptable option in selected cases.

Recently, an extended lymph node dissection comprising not only the obturator fossa, but also the external and the internal iliac area with the presacral nodes has been advocated (54,55) but this approach was not analyzed in a prospective randomized fashion. Nevertheless, the limited value of a lymph node dissection as a staging procedure only without any therapeutic benefit is nowadays more and more challenged.

### 8.7 Results of radical prostatectomy

The results achieved in a number of studies involving radical prostatectomy are shown in Table 6.

<table>
<thead>
<tr>
<th>Study</th>
<th>No. of patients</th>
<th>Mean follow-up (months)</th>
<th>5-year PSA-free survival (%)</th>
<th>10-year PSA-free survival (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Han, et al., 2001 (56)</td>
<td>2404*</td>
<td>75</td>
<td>84</td>
<td>74</td>
</tr>
<tr>
<td>Catalona &amp; Smith, 1994 (57)</td>
<td>925</td>
<td>28</td>
<td>78</td>
<td>65</td>
</tr>
<tr>
<td>Hull, et al. 2002 (58)</td>
<td>1000</td>
<td>53</td>
<td>-</td>
<td>75</td>
</tr>
<tr>
<td>Trapasso, et al., 1994 (59)</td>
<td>601</td>
<td>34</td>
<td>69</td>
<td>47</td>
</tr>
<tr>
<td>Zincke, et al., 1994 (60)</td>
<td>3170</td>
<td>60</td>
<td>70</td>
<td>52</td>
</tr>
</tbody>
</table>

* 15-year, 66%.
8.8 **Neoadjuvant hormonal therapy and radical prostatectomy**
Generally, neoadjuvant or up-front therapy is defined as therapy given prior to definitive local curative treatment (e.g. surgery or radiation therapy). As CaP is an androgen-dependent tumour, neoadjuvant hormonal therapy (NHT) is an appealing concept. Attempts to decrease the size of the prostate before radical prostatectomy were first reported by Vallett as early as 1944 (61).

In several studies of NHT in clinical stage T2 and T3 cancer, a decreased prostate volume and a lowering of the serum PSA levels have been reported after hormonal manipulation (62, 63). However, these trials were not randomized, there was no standard treatment protocol and the lengths of NHT varied considerably.

Five prospective, randomized studies have shown a decrease in positive surgical margin rates, with the use of a short-term (6 weeks-4 months) course of NHT (64-69). Follow-up of these randomized trials has indicated that this has not resulted in any difference in PSA-free failure after 3-5 years of follow-up (70-73). Since none of these studies was powered to study overall survival, the impact of NHT on overall survival remains unclear.

When surgical technique was considered, it was noted that surgery tended to be more difficult in pre-treated patients (66,74), but that the duration of radical prostatectomy, blood loss and number of transfusions were similar in NHT-treated patients and controls (65,66,74). The expectation had been that a longer duration of NHT could improve the PSA-free survival but a well designed randomized trial was unable to demonstrate any advantage of an eight-month over a 3-month preoperative hormonal treatment (75).

With these results in mind, NHT cannot be recommended for routine clinical use prior to radical prostatectomy. Also the optimistic results of a Southwest Oncology Group (SWOG) study on 4 months of NHT before radical prostatectomy for extensive T3 (and even T4) disease can only be appreciated, but need to be re-established by a randomized controlled study (47). Further studies on the duration, the type of androgen ablation or even early other medical treatments are needed in order to define its role in the treatment of localized or locally advanced CaP (76).

8.9 **Complications and functional outcome**
The postoperative complications of radical prostatectomy are listed in Table 7. The mortality rate is 0-1.5% (77), urinary fistulas are seen in 1.2-4% of patients (78) and urinary incontinence that persists after 1 year in 7.7% (79). In men undergoing prostatectomy, the rates of postoperative and late urinary complications are significantly reduced if the procedure is performed in a high-volume hospital and by a surgeon who performs a large number of such procedures (80-82).

Erectile dysfunction used to occur in nearly all patients, but nerve-sparing techniques can be applied in early-stage disease (83). Patients who benefit from nerve-sparing radical prostatectomy might have a higher chance of local disease recurrence and should therefore be carefully selected. Algorithms have been proposed to decrease the positive margin rate after nerve-sparing radical prostatectomy (84). Patients with poorly differentiated tumours, apical tumour extension and an intra-operatively palpable tumour are not suitable candidates for a nerve-sparing approach (85-87). Also unilateral nerve-sparing operations can be safely proposed (88). The early administration of intracavernous injection therapy could improve the definitive potency rates (89) and the significance of sural nerve transplant needs further multicentre study (90).

**Table 7: Complications of radical prostatectomy**

<table>
<thead>
<tr>
<th>Complication</th>
<th>Incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peri-operative death</td>
<td>0.0-2.1</td>
</tr>
<tr>
<td>Major bleeding</td>
<td>1.0-11.5</td>
</tr>
<tr>
<td>Rectal injury</td>
<td>0.0-5.4</td>
</tr>
<tr>
<td>Deep venous thrombosis</td>
<td>0.0-8.3</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>0.8-7.7</td>
</tr>
<tr>
<td>Lymphocele</td>
<td>1.0-3.0</td>
</tr>
<tr>
<td>Urine leak, fistula</td>
<td>0.3-15.4</td>
</tr>
<tr>
<td>Slight stress incontinence</td>
<td>4.0-50.0</td>
</tr>
<tr>
<td>Severe stress incontinence</td>
<td>0.0-15.4</td>
</tr>
<tr>
<td>Impotence</td>
<td>29.0-100.0</td>
</tr>
<tr>
<td>Bladder neck obstruction</td>
<td>0.5-14.6</td>
</tr>
<tr>
<td>Ureteral obstruction</td>
<td>0.0-0.7</td>
</tr>
<tr>
<td>Urethral stricture</td>
<td>2.0-9.0</td>
</tr>
</tbody>
</table>
8.10 SUMMARY OF RADICAL PROSTATECTOMY

8.10.1 Indications

- In patients with stage T1b-T2, Nx-N0, M0 disease and a life expectancy >10 years (level of evidence: 1b)

8.10.2 Optional

- Patients with a long life expectancy and stage T1a disease (level of evidence: 3)
- Patients with stage T3a disease, a Gleason score of > 8 and a PSA of < 20 ng/mL

8.10.3 Comments

- Short-term (3 months) neoadjuvant therapy with gonadotrophin releasing-hormone analogues is not recommended in the treatment of stage T1-T2 disease (level of evidence: 1a)
- Nerve-sparing surgery may be attempted in pre-operatively potent patients with low risk for extracapsular disease (T1c, Gleason score < 7 and PSA < 10 ng/mL or see Partin tables/nomograms) (level of evidence: 3)
- Unilateral nerve sparing procedures is an option in stage T2a disease (level of evidence: 4)
- The role of radical prostatectomy in patients with high-risk features, lymph-node involvement (stage N1 disease) or as a part of a planned multimodality treatment (with long-term hormonal and/or adjuvant radiation therapy), has not been evaluated (level of evidence: 4).

8.11 REFERENCES


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9 TREATMENT: DEFINITIVE RADIATION THERAPY

9.1 Introduction
There are no randomized studies that compare radical prostatectomy with either external beam therapy or brachytherapy for localized CaP, but the National Institutes of Health (NIH) consensus set up in 1988 (1) remains available: external irradiation offers the same long-term survival results as surgery; moreover, external irradiation provides a quality of life at least as good as that provided by surgery (2). In Europe, the 1990s saw the introduction of three-dimensional conformal radiotherapy (3D-CRT) and a growing interest in transperineal brachytherapy. At the onset of the third millennium, intensity modulated radiotherapy (IMRT), an optimized form of 3D-CRT, is gradually gaining ground in centres of excellence.

After the appropriate assessment of tumour extension, the choice of treatment must be made based on a multidisciplinary approach, taking into account the 2002 TNM classification, Gleason score, baseline PSA, age of the patient, comorbidity, life expectancy and quality of life. Obtaining a patient’s consent is essential after providing exhaustive information regarding diagnosis, the therapeutic modalities and morbidity.

Additional information on the various aspects of radiotherapy in the treatment of CaP is made available in a newly published extensive overview (3).

9.2 Technical aspects
Anatomical data, acquired by scanning the patient in a treatment position, are transferred to the 3D treatment planning system where the clinical target volume is visualized, following which a (surrounding) safety margin is added. At the time of irradiation, a multileaf collimator, automatically and continually, adapts to the contours of the target volume seen by each beam. The real-time verification of the irradiation field by means of portal imaging allows for comparison of the treated and simulated fields and correction of deviations where displacement is more than 5 mm. 3D-CRT is a high-precision technique which improves local control through dose escalation, without increasing the risk of morbidity. IMRT is possible with linear accelerators equipped with the latest multileaf collimators and specific software. The movement of the leaves during the course of the irradiation allows for the adaptation of the dose to be delivered within the treatment field and provides concave isodose curves.

Whatever the techniques and their sophistication, quality assurance plays a major role in the management of radiotherapy, due to the involvement of physicians, physicists, dosimetrists, radiographers, radiologists and computer scientists.

9.3 Localized CaP T1-2c N0, M0
9.3.1 T1a-T2a N0, M0 and Gleason score ≤ 6 and PSA < 10 ng/mL (low-risk group)
For external radiotherapy up to 70-72 Gy is recommended as it offers the same results as dose escalation (4).

9.3.2 T2b or PSA 10-20ng/mL, or Gleason score 7 (intermediate-risk group)
Many series have shown a significant impact of dose escalation on 5-year survival without biochemical relapse for patients classified as cT1c-T3, with a dose ranging from 76 to 81 Gy with no grade 3 or 4 late toxicity (4-6). This is the reason why intermediate-risk group patients may benefit from dose escalation, as shown by two randomized trials. The MD Anderson Cancer Centre randomized study compared 78 Gy 3D-CRT to a 70 Gy
conventional radiotherapy including 305 stage T1-3 patients with a pre-treatment PSA level of more than 10 ng/mL (median follow-up of 40 months). A significantly higher 5-year free-from-failure rate was found in 75% of the patients who received 78 Gy vs 48% of those who received 70 Gy (p=0.01) (7). This study has been confirmed by the PROG 95-09 interim analysis that evaluated 393 T1b-T2b patients – 75% of which with Gleason score of 6 or less – and with a PSA < 15 ng/mL. Patients were randomized to receive an initial boost to the prostate alone using conformal protons of either 19.8 or 28.8 GyE, then 50.4 Gy to a larger volume. With a median follow-up of 4 years, there was a significant decrease of the 5-year biochemical failure rate (p=0.00001) in favour of the patients assigned to the higher dose (79.2 GyE) versus those receiving a conventional dose (70.2 GyE) (8). In daily practice, although a consensus has not been reached yet concerning the level of the dose escalation, 78 Gy seems to represent a good compromise.

9.3.3 T2c, or Gleason score > 7 or PSA > 20 ng/mL (high-risk group)
External irradiation with dose escalation improves 5-year biochemical disease-free survival (7) but seems insufficient to cover the risk of relapse outside the pelvis. Many studies aim to evaluate the dose escalation with or without adjuvant hormonal therapy:

i) The MRC with neoadjuvant hormonal therapy comparing conventional radiotherapy of 64 Gy to high-dose (74 Gy) radical conformal radiotherapy (9)

ii) The Fédération Nationale des Centres de Lutte Contre le Cancer (FNCLCC) comparing 70 to 80 Gy without hormonal therapy (10)

iii) The European Organization for Research and Treatment of Cancer (EORTC) with dose stratification (70, 74 and 78 Gy) with or without neoadjuvant and concomitant hormonal therapy (11).

A prospective randomized trial, which included 206 patients with a PSA of at least 10 ng/mL (maximum 40 ng/mL), a Gleason score of at least 7 (range 5-10), or radiographic evidence of extra-prostatic disease, compared 3D-CRT alone or in combination with 6 months of androgen deprivation therapy (ADT). After a median follow-up of 4.5 years, patients randomized to receive 3D-CRT plus ADT had a significantly higher survival rate (p=0.04), lower CaP-specific mortality rate (p=0.02), and higher survival rate free of salvage ADT (p=0.002) (12).

9.3.4 Prophylactic irradiation of pelvic lymph nodes in intermediate- or high-risk localized CaP
Invasion of the pelvic lymph nodes is a poor prognostic indicator (13). Randomized trials held in the 1970s and 1980s failed to show that patients benefited from prophylactic irradiation of the pelvic lymph nodes in high-risk cases. The Radiation Therapy Oncology Group (RTOG) randomized study (1978-1983) with 484 patients T1b-T2 demonstrated that irradiation of the pelvic lymph node chains neither improved the rate of local recurrence nor survival (14), with results similar to those of a Stanford study (1970-1986) of only 91 patients (15). Nowadays, due to individual screening, comprehensive clinical work-up and new imaging modalities, the risk of pelvic lymph node invasion, may be assessed by Partin’s tables (16) or the Roach formula (17). The Roach formula estimates the risk of pelvic lymph node involvement higher than 15%: positive lymph node = 2/3 PSA + (GS-6) x 10. Another modality used is to perform selective sampling at coelioscopy (18) or mini-laparotomy and to decide upon treatment based on these findings.

9.4 Innovative techniques
9.4.1 Intensity modulated radiotherapy (IMRT)
IMRT enables radiation oncologists to homogeneously increase the doses up to 80 Gy within the target volume, while respecting the threshold doses in organs at risk. The Memorial Sloan-Kettering Cancer Centre have the largest experience with this technique, reporting on 772 patients treated between 1996 and 2001 with doses ranging from 81 to 86.4 Gy using an inverse planning approach. With a median follow-up time of 24 months (6-60 months), the 3-year actuarial likelihood of ≥ late grade 2 rectal toxicity was 4%; the 3-year actuarial likelihood of ≥ grade 2 urinary toxicity was 15%; and the 3-year actuarial PSA relapse-free survival rates for favourable-, intermediate- and unfavourable-risk group patients were 92%, 86% and 81% respectively (19). The use of IMRT is opening the way to hypofractionated treatment, with a shorter duration for the overall treatment time, by delivering 70 Gy in 28 fractions over 5.5 weeks, with 2.5 Gy per fraction (20).

9.4.2 Transperineal brachytherapy
Transperineal brachytherapy is a safe and efficient technique, which generally requires less than 2 days of hospitalization. There is a consensus on the following eligibility criteria: stage cT1b- T2a NO, M0, a Gleason score ≤ 6 assessed on a sufficient number of random biopsies, an initial PSA level of ≤ 10 ng/mL, a prostate volume of ≤ 50 cm³ and a good International Prostatic Symptom Score (IPSS) (21).

In 1983, Holm described the transperineal method with endorectal sonography in which the patient is positioned in a dorsal decubitus gynaecological position (22). Implantation is undertaken under general anaesthesia or spinal block and requires a learning curve from the whole team: the surgeon for the delineation
of the prostate and the placement of the needles, the physicist for real-time dosimetry, and the radiation oncologist for source loading. The sonography probe introduced into the rectum is fixed in a stable position. In cases of permanent implants, iodine-125 in granule form is the radio-element of reference; palladium-103 may be used for less differentiated tumours with high doubling time. The dose delivered to the planning target volume is in the order of 160 Gy for iodine-125 and of 120 Gy for palladium-103. A Gleason score of 7 still remains a “grey zone”, but patients with GS 4+3 show no difference in outcome (23). In cases of intermediate- or high-risk localized CaP, the combination with external irradiation (24) or neoadjuvant hormonal treatment (25) may be considered, but the potential positive impact of these treatments needs to be assessed with randomized trials. Non-permanent transperineal interstitial prostate brachytherapy using a high-dose rate iridium-192 stepping source and a remote afterloading technique can be applied with a total dose of 12 to 20 Gy in 2 to 4 fractions combined with fractionated external radiotherapy of 45 Gy (26).

For T1-2 N0 M0, the 5-year biochemical failure rates are similar for permanent seed implantation, high-dose (> 72 Gy) external radiation, combination seed/external irradiation, and radical prostatectomy. These were the results from a study including 2,991 patients diagnosed with T1-2 consecutive localized CaP treated between 1990 and 1998 at the Cleveland Clinic Foundation and Memorial Sloan-Kettering Cancer Centre with a minimum 1-year follow-up (27).

9.5 Late toxicity
Patients have to be informed about the potential late genitourinary or intestinal toxicity that may occur, as well as the impact of irradiation on erectile function. Late toxicity was analyzed in the prospective EORTC randomized trial 22863 (1987-1995) (28), where 90% of the patients included were diagnosed with T3-4. The specified dose was 70 Gy; 377 patients (91%) out of 415 enrolled were evaluable for long-term toxicity, graded according to a modified RTOG scale. There were 86 (22.8%) patients with grade ≥ 2 urinary or intestinal complications or leg oedema; of these 86 patients, 72 had grade 2 (moderate) toxicity, 10 had grade 3 (severe) toxicity, and 4 died due to grade 4 (fatal) toxicity. Although 4 (1%) late treatment-related deaths occurred, long-term toxicity was limited, with less than 5% grade 3 or 4 late complications being reported (Table 8). These data can be used as a baseline for comparison with irradiation techniques currently in use, such as 3D-CRT or IMRT. Radiotherapy affects erectile function to a lesser degree than surgery according to retrospective surveys of patients (2). A recent meta-analysis has shown that the 1-year rate of probability for maintaining erectile function after brachytherapy was 0.76, 0.60 after brachytherapy plus external irradiation, 0.55 after external irradiation, 0.34 after nerve sparing radical prostatectomy and 0.25 after standard radical prostatectomy. When studies with more than 2-years’ follow-up were selected (i.e. excluding brachytherapy), the rates became respectively: 0.60, 0.52, 0.25, 0.25, with a greater spread between the radiation techniques and the surgical approaches (29).

Table 8: Incidence of late toxicity by RTOG grade (from EORTC trial 22863)

<table>
<thead>
<tr>
<th>TOXICITY</th>
<th>Grade 2</th>
<th>Grade 3</th>
<th>Grade 4</th>
<th>Any significant toxicity (≥ grade 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cystitis</td>
<td>18 (4.7)</td>
<td>2 (0.5)</td>
<td>0 (0)</td>
<td>20 (5.3)</td>
</tr>
<tr>
<td>Hematuria</td>
<td>18 (4.7)</td>
<td>0</td>
<td>0</td>
<td>18 (4.7)</td>
</tr>
<tr>
<td>Urinary stricture</td>
<td>18 (4.7)</td>
<td>5 (1.3)</td>
<td>4 (1)</td>
<td>27 (7.1)</td>
</tr>
<tr>
<td>Urinary incontinence</td>
<td>18 (4.7)</td>
<td>2 (0.5)</td>
<td>0 (0)</td>
<td>20 (5.3)</td>
</tr>
<tr>
<td>Overall GU Toxicity</td>
<td>47 (12.4)</td>
<td>9 (2.3)</td>
<td>4 (1)**</td>
<td>60 (15.9)</td>
</tr>
<tr>
<td>Proctitis</td>
<td>31 (8.2)</td>
<td>0</td>
<td>0</td>
<td>31 (8.2)</td>
</tr>
<tr>
<td>Chronic diarrhoea</td>
<td>14 (3.7)</td>
<td>0</td>
<td>0</td>
<td>14 (3.7)</td>
</tr>
<tr>
<td>Small bowel obstruction</td>
<td>1 (0.2)</td>
<td>1 (0.2)</td>
<td>0</td>
<td>2 (0.5)</td>
</tr>
<tr>
<td>Overall GI Toxicity</td>
<td>36 (9.5)</td>
<td>1 (0.2)</td>
<td>0</td>
<td>37 (9.8)</td>
</tr>
<tr>
<td>Leg Oedema</td>
<td>6 (1.5)</td>
<td>0</td>
<td>0</td>
<td>6 (1.5)</td>
</tr>
<tr>
<td>Overall Toxicity*</td>
<td>72 (19)</td>
<td>10 (2.7)</td>
<td>4 (1)</td>
<td>86 (22.8)</td>
</tr>
</tbody>
</table>

(GU&GI toxicity and leg oedema)

* Most of the patients had more than one type of toxicity thus the total at the bottom of the table do not result from simple addition.

** 2 of the grade 4 patients were irradiated with CO 60

Note: There was not any other significant (≥ grade 2) toxicity among patients irradiated with CO 60 (n=15) except for the 2 grade 4 GU toxicity (stated above) and only 1 patient with grade 2 GI toxicity.
9.6 Immediate postoperative external irradiation for pathological tumour stage T3 N0, M0
Extracapsular invasion (pT3) is burdened with a risk of local recurrence, which can reach up to 30% (30); in multifactorial analysis, the predictors of biochemical relapse are: PSA level (p=0.005); Gleason score of the surgical specimen (p=0.002); and positive surgical margins (p < 0.001) (31). Only one prospective randomized trial has assessed the role of immediate postoperative radiotherapy; EORTC study 22911 compared immediate postoperative radiotherapy (60 Gy) to radiotherapy delayed until local recurrence (70 Gy) in patients classified as pT3 pN0 after retro-pubic radical prostatectomy. Immediate postoperative radiotherapy proved to be well tolerated with a risk of grade 3-4 urinary toxicity of under 3.5% (32), without significant difference regarding incontinence and/or stricture of anastomosis (33). The study concludes that immediate postoperative radiotherapy after surgery significantly improves 5-year clinical or biological survival: 72.2% vs 51.8% p < 0.0001 (34).

Consequently, for patients classified as T1-2 N0 (or T3 N0 with selected prognostic factors), pT3 pN0 with high risk of local failure after radical prostatectomy due to rupture of the capsule, positive margins and/or invasion of the seminal vesicles, presenting with a PSA of < 0.1 ng/mL one month after surgery, the following may be recommended:
• Immediate radiotherapy upon recovery of urinary function.
• Or, clinical and biological monitoring followed by salvage radiotherapy, when the PSA exceeds 0.5 ng/mL (35); 1.0 ng/mL seems to be a breakpoint, above which the likelihood of local control is significantly reduced (36).

9.7 Locally advanced CaP: T3-4 N0, M0, T1-4 N1 M0
The incidence of locally advanced CaP declined as a result of individual or mass screening. Pelvic lymph node irradiation is optional for N0 patients, due to the likelihood of infraclinical disease and N1 patients (inter-iliac nodes). Results of radiotherapy alone are dismal (37). This is why, because of the hormone dependence of CaP (38), ADT has been combined with external irradiation with the dual objectives of:
• Reducing the risk of distant metastases by potentially sterilizing micrometastases already present at the moment of diagnosis.
• Decreasing the risk of non-sterilization and/or local recurrence as a source of secondary metastases (39) through the effect of radiation-induced apoptosis (40,41).
Numerous randomized trials have assessed the value of this combination.

9.7.1 Neoadjuvant hormonal therapy
The RTOG study 86-10 included 471 patients with stage T2-4N0-X M0. ADT was administered 2 months before irradiation and during irradiation, or in the case of relapse in the control arm. 32% of patients were diagnosed as T2, 70% as T3-4 and 91% N0. The hormone treatment consisted of oral eulexine, 250 mg, 3 times daily and goserelin acetate (Zoladex) 3.6 mg every 4 weeks by subcutaneous injection. The pelvic target volume received 45 Gy and the prostatic target volume received 20-25 Gy. At 8 years, ADT was associated with an improvement in local control (42% vs 30%, p=0.016), disease-free survival (33% vs 21%, p=0.004) and biochemical disease-free survival (PSA < 1.5 ng/mL, 24% vs 10% (p < 0.0001)). In patients with Gleason score 2-6, there was a significant improvement in survival: 70% vs 52% (p=0.015) (42).

9.7.2 Concomitant and adjuvant hormonal therapy
The EORTC study 22863 recruited 415 patients diagnosed with T1-2 grade 3 WHO (World Health Organization), T3-4 N0, M0 and compared radiotherapy with adjuvant ADT to radiotherapy alone. ADT was allowed in cases of relapse. 82% of patients were diagnosed as T3, 10% as T4 and 89% as N0. The hormone treatment consisted of oral cyproterone acetate, 50 mg 3 times daily for 1 month, beginning 1 week before the start of radiotherapy and subcutaneous injection of goserelin acetate 3.6 mg every 4 weeks for 3 years, starting on the first day of radiotherapy.
The pelvic target volume received was 50 Gy and the prostatic target volume was 20 Gy. With a median follow-up of 66 months, combination therapy compared with radiotherapy alone was significantly better for both survival (78% vs 62%, p=0.001) and survival without clinical relapse (78% vs 40%, p < 0.001) (43). The 5-year cumulative incidence of locoregional failure was 1.7% vs 16.4% in the radiotherapy alone arm (p < 0.0001) and survival without clinical or biological failure (nadir of 1.5 ng/ml) was 81% for the combined treatment arm vs 43% in the radiotherapy alone arm (p < 0.001).

9.7.3 Adjuvant hormonal therapy
The RTOG study 85-31 recruited 977 patients diagnosed with T3-4 N0-1 M0, or pT3 after radical prostatectomy. ADT was begun in the last week of irradiation and continued up to relapse (Group I) or started at recurrence (Group II). 15% of patients in Group I and 29% in Group II had undergone radical prostatectomy, while 14% of patients in Group I and 26% in Group II were pN1. Goserelin acetate 3.6 mg was administered
every 4 weeks. The pelvis received 45 Gy and the prostatic bed received 20-25 Gy. Patients diagnosed with stage pT3 received 60-65 Gy. With a median follow-up time of 7.3 years, a statistical significance was reached for 5-year and 10-year overall survival in favour of the adjuvant hormonal therapy arm, 76 vs 71% and 53 vs 38% respectively (44). In this study, 95 of the 173 pN1 patients who received pelvic radiotherapy with immediate hormonal therapy had a significantly better survival rate without biochemical relapse at 5 years (PSA < 1.5 ng/mL) than those in the arm with delayed hormonal therapy (p=0.0001) (45).

The National Cancer Institute (NCI) Canada study, including patients diagnosed with stage cT3-4 N0 M0, compared complete androgen blockade (CAB) (goserelin acetate 3.6 mg subcutaneous every 4 weeks with flutamide 750 mg/day) alone and in combination with radiation 65-69 Gy (46). The results are awaited.

9.7.4 Neoadjuvant, concomitant and adjuvant hormonal therapy
The RTOG 92-02 trial closed in 1995 after accruing 1,554 patients. Statistically significant improvements were observed in bNED (actuarial biochemical freedom of disease) control, distant metastatic failure, local control, and disease-free survival for patients receiving long-term ADT (before, during, and 2 years after radiotherapy) compared with short-term treatment (2 months before, and during, radiotherapy). With a median follow-up of 5.8 years, the long-term androgen deprivation (ablation) (LTAD) arm showed significant improvement in all efficacy end-points except 5-year overall survival, 80% vs 76.5% (p=0.73), compared with the short-term androgen deprivation (ablation) (STAD). In a subset of patients, who were not part of the original study design, with Gleason score 8-10 tumours, after 5 years the LTAD arm showed significantly better overall survival: 81% vs 70.7%, (p=0.04) (47).

9.8 SUMMARY OF DEFINITIVE RADIATION THERAPY
1. In localized CaP T1c-T2c N0 M0, 3D-CRT with or without IMRT, is recommended, even for young patients who refuse surgical intervention. There is fairly strong evidence that intermediate-risk patients benefit from dose escalation (level of evidence: 2). For patients in the high-risk group short-term ADT prior to, and during, radiotherapy may result in increased overall survival (level of evidence: 2a)
2. Transperineal interstitial brachytherapy with permanent implants may be proposed to patients cT1-T2a, Gleason score < 7 (or 3+4), PSA ≤ 10 ng/mL, prostate volume ≤ 50 mL, without a previous TURP and with a good IPSS (level of evidence: 2b)
3. Immediate postoperative external irradiation after radical prostatectomy for patients with pathological tumour stage T3 N0 M0 prolongs biochemical and clinical disease-free survival (level of evidence: 2a). An alternative option is to give radiation at the time of biochemical failure but before PSA reaches above 1-1.5 ng/mL (level of evidence: 3)
4. In locally advanced CaP, overall survival is improved by concomitant and adjuvant hormonal therapy (with a total duration of 2 to 3 years) with external irradiation (level of evidence: 1). For a subset of patients, T2c-T3 N0-x with Gleason score 2-6, short-term ADT before, and during, radiotherapy may favourably influence overall survival (level of evidence: 1b).

9.9 REFERENCES


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10 EXPERIMENTAL LOCAL TREATMENT OF PROSTATE CANCER

10.1 Background
Besides radical prostatectomy, external beam radiation and/or brachytherapy, cryosurgery of the prostate (CSAP) and high-intensity focused ultrasound (HIFU) have emerged as alternative therapeutic options in patients with clinically localized CaP (1-4). Whereas HIFU is still considered to be an experimental treatment, CSAP has been recognized as a true therapeutic alternative as recommended by the guidelines of the American Urological Association. Both techniques have been developed as minimally invasive procedures potentially resulting in the same therapeutic efficacy as the established surgical and non-surgical options associated with reduced therapy-associated morbidity.

10.2 Cryosurgery of the prostate (CSAP)
Cryosurgery uses freezing techniques to induce cell death by: (1) dehydration resulting in protein denaturation; (2) direct rupture of cellular membranes by ice crystals; (3) vascular stasis and microthrombi resulting in stagnation of the microcirculation with consequent ischemia; and (4) apoptosis (1-4). Freezing of the prostate is ensured after placement of 12-15 17G cryoneedles under TRUS guidance, placement of thermosensors at the level of the external sphincter and the bladder neck and insertion of a urethral warmer. Two freeze-thaw cycles are used under TRUS guidance resulting in a temperature of -40° Celsius in the mid gland and at the neurovascular bundle.

10.2.1 Indication for CSAP
Patients ideally suitable for CSAP are those who have organ-confined CaP and those identified to have minimal extension beyond the prostate (1-3). Prostate size should be ≤ 40mL; prostate glands > 40mL should be hormonally downsized in order to prevent technical difficulties in placing cryoprobes under the pubic arch. PSA serum levels should be < 20ng/mL and the biopsy Gleason score should be < 7. Since there are no, or only very few, data on the long-term outcome in terms of cancer control at 10 and 15 years, patients with a life expectancy > 10 years have to be informed accordingly.
10.2.2 Results of modern cryosurgery for CaP

When comparing treatment modalities, it is important to keep in mind that in modern radical prostatectomy series of patients with clinically organ-confined CaP, the risk of dying from CaP 10 years after surgery is as low as 2.4% (5).

Therapeutic results have improved over time with improved techniques in terms of gas-driven probes and transperineal probe placement as used in third-generation cryosurgery (6-12). Objective assessment of PSA outcome cannot easily be performed since some institutions used PSA values < 0.1 ng/mL as an indicator of therapeutic success, whereas others used the American Society of Therapeutic Radiology and Oncology (ASTRO) criteria with three consecutive PSA increases. With regard to second-generation CSAP, if a PSA nadir < 0.5 ng/mL is used, biochemical free survival at 5 years is 60% and 36% for low-risk and high-risk patients, respectively (6,7). The 7-year biochemical free survival, however, is 92% if ASTRO criteria are used. Long et al. (6) retrospectively analyzed the multicentre pooled CSAP results of 975 patients who were stratified into three risk groups. Using PSA thresholds of 1.0 ng/mL and < 0.5 ng/mL at a mean follow-up of 24 months, the 5-year actuarial biochemical progression-free rate was 76% and 60%, respectively for the low-risk group, 71% and 45%, respectively for the intermediate-risk group and 61% and 36%, respectively for the high-risk group.

With regard to third-generation cryosurgery, clinical follow-up is short with only 110/175 (63%) patients having a PSA follow-up at 12 months (6-12). 80 (73%) patients remain with a PSA nadir < 0.4 ng/mL and 42/65 (76%) low-risk patients remain free from biochemical progression using the 0.4 ng/mL cut-off. Longer follow-up was reported by Bahn, et al. (9) analyzing the therapeutic results of 590 patients undergoing CSAP for clinically localized and locally advanced CaP. Using a PSA cut-off level of < 0.5 ng/mL, the 7-year biochemical-free survival for low-, medium- and high-risk groups were 61%, 68% and 61%, respectively.

Nerve-sparing cryosurgery, as reported recently (13), must still be considered to be an experimental therapeutic option. Nerve-sparing surgery was performed in 9 patients with unilateral CaP confirmed on repeated biopsies, with CSAP performed on the side of the positive biopsy whereas the negative biopsy side was spared from freezing.

10.2.3 Complications of CSAP for primary treatment of CaP

Erectile dysfunction occurs in about 80% of the patients and remains a consistent complication of the CSAP procedure, independent of the system generation used. The complication rates described with the third-generation cryosurgery include tissue sloughing in about 3%, incontinence in 4.4%, pelvic pain in 1.4% and urinary retention in about 2% (6-12). The development of fistula is usually rare with < 0.2% in modern series. About 5% of all patients require TURP for subvesical obstruction.

Quality of life and sexuality following CSAP have been investigated in a clinical phase II trial recruiting 75 men (14). Quality-of-life analysis by the prostate-specific FACT-P questionnaire revealed that most subscales had returned to pre-treatment levels by 12 months after CSAP. Furthermore, no significant changes could be determined when comparing the 36-month with the 12-month data. With regard to sexuality, 37% of the men were able to have intercourse at 3 years after CSAP.

10.2.4 Summary of CSAP

- Patients with low-risk CaP (PSA < 10 ng/mL, ≤ T2a, Gleason score ≤ 6) or intermediate-risk CaP (PSA > 10 ng/mL or Gleason score ≥ 7 or stage ≥ 2b) represent potential candidates for CSAP.
- Prostate size should be < 40 mL at the time of therapy.
- Long-term results are lacking and 5-year biochemical progression-free rates are inferior to those achieved by radical prostatectomy in low-risk patients. Patients have to be informed accordingly.

10.3 High-intensity focused ultrasound (HIFU) of the prostate

HIFU consists of focused ultrasound waves emitted from a transducer inducing tissue damage by mechanical and thermal effects as well as by cavitation (15). The goal of HIFU is to heat malignant tissues above 65°C to destroy these tissues by coagulative necrosis. HIFU is performed under general or spinal anaesthesia with the patient in the lateral position; the procedure is time consuming with about 10g prostate tissue being treated in 1 hour.

10.3.1 Results of HIFU in CaP

As with CSAP, it appears to be difficult to interpret oncological outcome in patients undergoing HIFU since various PSA thresholds are defined and no international consensus exists on objective response criteria. Results of HIFU are limited with the outcome data of less than 1,000 CaP cases being published in the literature. In one of the studies (16), a significant decrease of pre-treatment PSA serum levels from 12 ng/mL to 2.4 ng/mL was observed. However, 50% of the 14 patients demonstrated positive prostate biopsies during follow-up. In another study (17), a complete response rate defined by PSA < 4 ng/mL and 6 negative biopsies could be achieved in 56% of the patients.
Summarizing the efficacy results of a European multicentre study comprising the data of 559 patients with mainly low- and intermediate-risk CaP, Thüroff et al. (17) reported on a negative biopsy rate of 87.2% in 288 men with a follow-up of at least 6 months. PSA nadir after 6 months follow-up could be determined in 212 patients and it was as high as 1.8 ng/mL. However, it could be demonstrated that the PSA nadir might be reached at 12 to 18 months following the initial procedure. Blana et al. reported 146 patients undergoing HIFU with a mean follow-up of 22.5 months (18). The mean PSA level at initiation of therapy was 7.6 ng/mL, the PSA nadir achieved after 3 months was 0.07 ng/mL. However, after 22 months the median PSA level was 0.15 ng/mL. Of 137 men available for analysis, 93.4% demonstrated a negative control biopsy.

10.3.2 Complications of HIFU
Urinary retention appears to be one of the most common side-effects of HIFU developing in basically all patients, with the mean interval of catheterization via a suprapubic tube varying between 12 and 35 days (16-18). Grade I and II urinary stress incontinence occurs in about 12% of the patients. Subsequent TURP or bladder neck incision to treat subvesical obstruction is common and sometimes even performed at the time of HIFU. Postoperative impotence will occur in approximately 55% to 70% of the patients.

10.4 Radiofrequency interstitial tumour ablation (RITA)
RITA is a recently developed minimally invasive therapeutic option delivering radiofrequency energy via a needle electrode placed inside the prostate and resulting in coagulative necrosis by heating the tissue up to 100°C. Clinical application so far has been limited to two small studies demonstrating the feasibility and safety of the procedure (19,20). However, there are reliable data with regard to oncological control of CaP.

10.5 SUMMARY OF EXPERIMENTAL THERAPEUTIC OPTIONS TO TREAT CLINICALLY LOCALIZED CAP
1. CSAP has evolved from an investigational therapy to a possible alternative to treat CaP in patients unfit for surgery or in those with a life expectancy < 10 years (grade C recommendation)
2. All other minimally invasive treatment options, such as HIFU, RITA, microwaves and electrosurgery, are still experimental or investigational. For all of these procedures, a longer follow-up is mandatory to assess their true role in the management of CaP (grade C recommendation).

10.6 REFERENCES


11 HORMONAL THERAPY

11.1 Introduction
In 1941, Huggins and Hodges assessed the favourable effect of surgical castration and oestrogen administration on the progression of metastatic CaP, demonstrating for the first time the responsiveness of CaP to androgen deprivation (1,2).

Since their pivotal studies, androgen-suppressing strategies have become the mainstay for the management of advanced CaP, but recent years show an evolution towards increasing hormonal treatment of younger men with earlier (i.e., non-metastatic) stages of disease or recurrent disease after definitive treatment; either as primary single-agent therapy or as a part of a multimodality approach (3).

Even if hormonal treatment effectively palliates the symptoms of advanced disease, there is no conclusive evidence at present that it can extend life.

11.2 Basics of hormonal control of the prostate
Prostate cells are physiologically dependent on androgens to stimulate growth, function and proliferation. Testosterone, although not tumourigenic, is essential for the growth and perpetuation of tumour cells (4). The testes are the source of the vast majority of the androgens with only 5 to 10% – androstenedione, dihydroepiandrosterone and dihydroepiandrosterone sulphate – deriving from adrenal biosynthesis.

Testosterone secretion is regulated by the hypothalamic-pituitary-gonadal axis. The hypothalamic luteinizing hormone-releasing hormone (LHRH) stimulates the anterior pituitary gland to release luteinizing hormone (LH) and follicle-stimulating hormone (FSH). LH stimulates the Leydig cells of the testes to secrete testosterone. Within the prostate cells, testosterone is converted by the enzyme 5-α-reductase into 5-α-dihydrotestosterone (DHT), which is an androgenic stimulant approximately 10 times more powerful than the parent molecule (5). Circulating testosterone is peripherally aromatized and converted into oestrogens, which, together with the circulating androgens, exert a negative feedback control on the hypothalamic LH secretion.

If prostate cells are deprived of androgenic stimulation, they undergo apoptosis (programmed cell death). Any treatment ultimately resulting in the suppression of androgen activity is referred to as androgen deprivation therapy (ADT).

11.3 Different types of hormonal therapy
Androgen deprivation can be achieved either by suppressing the secretion of testicular androgens by means of surgical or medical castration, or by inhibiting the action of the circulating androgens at the level of their receptor in prostate cells using competing compounds known as antiandrogens. Alternatively, these two modalities can be combined in order to achieve what is commonly known as complete (or maximal or total) androgen blockade (CAB).

11.3.1 Testosterone-lowering therapy (castration)
11.3.1.1 Bilateral orchietomy
Surgical castration is still considered the “gold standard” for ADT against which all other treatments are rated.

By removing the testicular source of androgens, a hypogonadal status with a considerable decline of testosterone concentrations is induced, though a very low level of testosterone (known as “castration level”) does persist. Bilateral orchietomy, either by means of total or subcapsular (i.e., with preservation of tunica albuginea and epididymis) technique, is a simple and virtually complication-free surgical procedure, which can easily be performed under local anaesthesia (6).

The main drawback of orchietomy is that it may have a negative psychological effect; some men consider it to be an unacceptable assault on their manhood.

In recent years, a decline in the utilization of bilateral orchietomy can be witnessed which can be attributed to effects of stage migration towards earlier disease and the introduction of equally effective pharmacological modalities of castration (7).
11.3.1.2 Oestrogens
The mechanism of action is multifold: down-regulation of LHRH secretion, androgen inactivation, direct suppression of Leydig cell function and direct cytotoxicity to the prostate epithelium (only in-vitro evidence) (8).

The most commonly used oestrogen is diethylstilboestrol (DES). In early studies by the Veterans Administration Co-operative Urological Research Group (VACURG) (9), oral DES at a dosage of 5 mg/day was tested, but the treatment was associated with high cardiovascular morbidity and mortality due to the first-pass hepatic metabolism with formation of thrombogenic metabolites. Accordingly, subsequent studies (10) tested lower oral dosages, namely 3 mg and 1 mg: both regimens provided a therapeutic efficacy comparable to that of bilateral orchiectomy, but the former was still associated with high cardiotoxicity. Although a 1 mg dose was associated with substantially less cardiovascular adverse events than the 5 mg dosage, the side-effects were still significantly increased compared to castration. Due to these concerns and the advent of LHRH agonists and antiandrogens, the use of DES had fallen out of favour until recently.

Renewed interest in oestrogens can be ascribed to three main reasons: firstly, as a response to the number of deleterious side-effects and high costs of long-term ADT with the nowadays widespread LHRH agonists: oestrogens suppress testosterone levels and do not seem to lead to bone loss and cognitive decline (11, level of evidence: 3); secondly, oestrogenic compounds (DES, DES-diphosphate, and the herbal supplement, PC SPES) have been shown to induce PSA-response rates as high as 86% in phase II trials with patients diagnosed with hormone-refractory prostate cancer (HRPC); thirdly, a new oestrogen receptor-beta (ER-b), possibly involved in prostate tumourigenesis, has been discovered (8).

Two different strategies have been used in an attempt to neutralize the cardiotoxicity, which is the main drawback of oestrogen therapy. These strategies are using the parenteral route of administration, which avoids hepatic first-pass metabolism, and the addition of cardiovascular protecting agents. The final analysis of the Scandinavian Prostatic Cancer Group Study 5 (a prospective randomized trial of more than 900 men with metastatic CaP that compared a parenteral oestrogen (polyestradiol phosphate) versus CAB [orchiectomy or LHRH agonist plus flutamide]) showed neither a significant difference in disease-specific and overall survival between the treatment arms nor a significant increase in cardiovascular mortality in the oestrogen arm, although the occurrence of non-fatal cardiovascular adverse events was considerably higher in this group (12). On the other hand, third recent, though small, phase II trials of patients with advanced CaP or HRPC evaluated the combination of DES, 1 or 3 mg/day, with either low-dose (1 mg/day) warfarin sodium or low-dose (75-100 mg/day) aspirin in the prevention of cardiovascular toxicity and found a persistent rate of thromboembolic complications (13-15).

In conclusion, DES is one of the classic forms of hormonal therapy. Although its efficacy was demonstrated many years ago and recently reconfirmed in a meta-analysis as comparable to that of bilateral orchiectomy (16, level of evidence: 1a), the significant cardiovascular side-effects, even at lower dosages, remain a concern. Further data are needed before oestrogens will be readmitted in clinical practice as a standard first-line treatment option.

11.3.1.3 Luteinizing hormone-releasing hormone (LHRH) agonists
Long-acting LHRH agonists (buserelin, goserelin, leuprorelin and triptorelin) have been used in advanced CaP for more than 15 years and are currently the predominant forms of ADT (3,17).

They are synthetic analogues of LHRH, generally delivered as depot injections on a 1-, 2- or 3-month basis, that interfere with the hypothalamic-pituitary-gonadal axis. They initially stimulate pituitary LHRH receptors, inducing a transient rise in LH and FSH release and consequently elevate testosterone production (known as “testosterone surge” or “flare up” phenomenon), which begins approximately within 2 or 3 days after the first injection and lasts through approximately the first week of therapy (18). Chronic exposure to LHRH agonists eventually results in down-regulation of LHRH-receptors, with subsequent suppression of pituitary LH and FSH secretion and testosterone production. The level of testosterone decreases to castration levels usually within 2 to 4 weeks (19,20). However, approximately 10% of patients treated with LHRH agonist fail to achieve castration levels (21).

In a recent meta-analysis evaluating single-therapy ADT for advanced CaP, LHRH agonists have shown comparable efficacy to orchiectomy and DES (16, level of evidence: 1a). In addition, although only based on an indirect comparison, all seemed equally effective (16, level of evidence: 3).

Today, LHRH agonists have become the “standard of care” in hormonal therapy because they avoid the physical and psychological discomfort associated with orchiectomy and lack the potential cardiotoxicity associated with DES. However, the main concerns associated with the administration of LHRH agonists are the potentially detrimental effects associated with the “flare phenomenon” in advanced disease, namely increased bone pain, acute bladder outlet obstruction, obstructive renal failure, spinal cord compression and fatal cardiovascular events due to hypercoagulation status. A recent review (22) addressing these issues concluded that clinical flare needs to be distinguished from the more common biochemical flare (i.e. increasing levels of PSA) and even from asymptomatic radiographic evidence of progression, and that patients at risk for clinical flare are overwhelmingly those with high-volume, symptomatic, bony disease, accounting for only 4-10% of M1
patients. Concomitant therapy with an antiandrogen definitely decreases the incidence of clinical relapse, but it does not completely remove the possibility of their occurrence. Based on pharmacokinetic considerations, it is recommended that administration of the antiandrogens should be started on the same day as the depot injection and treatment should be continued for a 2-week period. However, for patients with impending spinal cord compression, alternative strategies for immediately ablating testosterone levels must be considered, such as bilateral orchietomy or LHRH-antagonists.

11.3.1.4 LHRH antagonists
In contrast to the agonists, LHRH antagonists bind immediately and competitively to LHRH receptors in the pituitary gland. The effect is a rapid decrease in LH, FSH and testosterone levels without any flare. This seemingly more desirable mechanism of action has made LHRH antagonists very attractive since their introduction, but practical shortcomings have limited clinical studies. Indeed, many of these compounds have been associated with serious and life-threatening histamine-mediated side-effects and, until recently, no depot formulation was available.

Two recently published phase III randomized multicentre trials comparing the LHRH antagonist abarelix with the LHRH agonist leuprorelin acetate (23) and with CAB (24) in patients with metastatic or recurrent CaP showed no difference in achieving and maintaining castration levels of testosterone and in reducing serum PSA; the biochemical “flare up” phenomenon was not reported in the abarelix arms and the overall incidence of severe adverse events (including allergic reactions) was similar across all treatment groups. Data on survival endpoints and long-term safety are not yet available.

Abarelix has recently been licensed by the United States Food and Drug Administration for clinical use, but its use is restricted to those patients with metastatic and symptomatic CaP for whom no other treatment option is available (25).

11.3.2 Antiandrogens
Antiandrogens compete with testosterone and DHT for binding sites on their receptors in the prostate cell nucleus, thus promoting apoptosis and inhibiting CaP growth (26).

These orally administered compounds are classified according to their chemical structure as steroidal (e.g., cyproterone acetate (CPA), megestrol acetate and medroxyprogesterone acetate) and non-steroidal or pure (e.g., nifluramine, flutamide and bicalutamide). Both classes act as competitors of androgens at the receptor level, but while this is the sole action of non-steroidal antiandrogens, steroidal antiandrogens additionally have progestational properties with central inhibition of the pituitary gland. As a consequence non-steroidal antiandrogens do not lower testosterone levels, which remain normal or, conversely, slightly elevated.

11.3.2.1 Steroidal antiandrogens
These compounds are synthetic derivatives of hydroxyprogesterone. In addition to peripherally blocking androgen receptors, they have progestational properties and inhibit gonadotrophin (LH and FSH) release and suppress adrenal activity. At high doses megestrol acetate is cytotoxic.

Since steroidal antiandrogens lower testosterone levels, the main pharmacological side-effects are loss of libido and erectile dysfunction; gynaecomastia is quite rare. The non-pharmacological side-effects are cardiovascular toxicity (4-40% for CPA) and hepatotoxicity.

Cyproterone acetate
CPA was the first antiandrogen to be licensed and is the most widely used. There is only one randomized trial (27) comparing CPA to standard hormonal therapy (i.e. medical castration): patients in arm A (no contraindications to DES) were randomly assigned to CPA, goserelin or DES, while patients in arm B (contraindications to DES) were assigned to CPA or goserelin. In arm A, treatment with CPA was associated with significantly poorer median overall survival than goserelin only: adjusting for baseline characteristics did not account for this difference.

Two other studies on CPA monotherapy were performed, but one did not report survival data (28) and the other used non-standard treatment combination (DES and medroxyprogesterone acetate [29]). It is therefore difficult to draw any definite conclusions from these data about the relative efficacy of CPA and castration. Since no dose-finding studies of CPA monotherapy have been conducted, the most effective dose is still unknown. Although CPA has a relatively long half-life (30-40 hours), it is usually administered in two or three fractional doses of 100 mg each (30).

The only comparative study on antiandrogens as monotherapy was recently published by the European Organization for Research and Treatment of Cancer (EORTC) Protocol 30892 (a randomized trial of 310 patients comparing CPA versus flutamide in metastatic CaP), which showed no difference in cancer-specific and overall survival at a median follow-up of 8.6 years, though the study was underpowered (31) (level of evidence:1b).
Megesterol acetate and medroxyprogesterone acetate

Very limited information is available on these two compounds. Early studies with megestrol acetate demonstrated symptomatic and partial beneficial clinical response both in previously untreated metastatic CaP (32-34) and, to a lesser extent, in HRPC (35). No apparent dose-response correlation was shown to exist in a recent trial (36). The overall poor efficacy precluded megestrol acetate and medroxyprogesterone acetate from being recommended as a primary or second-line hormonal therapy option.

The only prospective randomized trial evaluating medroxyprogesterone acetate as primary therapy in advanced (M0-1) CaP is the EORTC 30761 study mentioned above (28), in which 236 patients were assigned to receive CPA, DES or medroxyprogesterone acetate: while no difference in cancer-specific and overall survival was evident between CPA and DES, treatment with medroxyprogesterone acetate had a less favourable course with a shorter survival time and time to progression than any of the other two drugs tested.

11.3.2.2 Non-steroidal antiandrogens

Non-steroidal antiandrogens have been promoted in monotherapy for quality of life (QoL) and compliance benefits over castration since they do not suppress testosterone secretion; it is claimed that libido, overall physical performance and bone mineral density are preserved (37).

Although no direct comparisons have been undertaken in a monotherapy setting, the three available drugs do not appear to differ in the severity of pharmacological side-effects, namely gynaecomastia, breast pain and hot flashes. However, there are differences in the non-pharmacological side-effects, with bicalutamide showing a more favourable safety and tolerability profile than nilutamide and flutamide (38).

Nilutamide

There are no comparative trials on nilutamide monotherapy with castration or with other antiandrogens (39). Only one non-comparative study was carried out, including 26 patients with M1 CaP who received nilutamide 100 mg three times daily. The results showed that as few as 38.5% of patients experienced an objective response; the median progression-free survival time was 9 months and the median overall survival was 23 months (40).

One large randomized controlled trial of 457 patients with M1 comparing orchiectomy plus nilutamide 300 mg/day vs orchiectomy plus placebo showed a significant benefit in cancer-specific and overall survival for the combined therapy (41). Recently nilutamide has been tested as second-line hormonal therapy in HPRC with encouraging results (42, 43). Non-pharmacological side-effects are visual disturbances (i.e., delayed adaptation to darkness), alcohol intolerance, nausea, hepatotoxicity and interstitial pneumonitis. Nilutamide is not licensed as a drug for monotherapy.

Flutamide

Flutamide was the first non-steroidal antiandrogen available for clinical use and has been studied as monotherapy for over 20 years, but no dose-finding studies against a currently accepted endpoint (e.g., PSA response) have been published. Flutamide is a pro-drug and the half-life of the active metabolite is 5 to 6 h, so it has to be administered three times daily to maintain therapeutic serum levels; the recommended daily dosage is 750 mg (30).

Early phase II trials demonstrated flutamide to be effective in the treatment of advanced CaP, albeit that the reported response rates cannot be correlated with currently recommended endpoints. The main advantage shown in these studies was the preservation of sexual function, which was maintained in up to 80% of patients with no pre-treatment erectile dysfunction (44-47). This rate has not been confirmed in the above mentioned EORTC trial 30892 (31), where as few as 20% of men treated with flutamide maintained sexual activity for up to 7 years.

Although several phase III studies have been conducted, results are often difficult to evaluate because of several drawbacks, such as the use of non-standard combinations, short-term follow-up and underpowering. Of these studies, only two phase III randomized trials comparing flutamide monotherapy to standard therapy (orchiectomy [48] and CAB [49]) for advanced CaP have reported survival data; both showed no significant difference in overall survival for flutamide or castration. Results are eagerly awaited from an ongoing Swedish study in which 700 patients with M1 CaP have been randomized to flutamide 250 mg three times daily or CAB (37).

The non-pharmacological side-effects are diarrhoea and hepatotoxicity (occasionally fatal).

Bicalutamide

Early reports with bicalutamide monotherapy related only to the 50 mg dosage, which was the one licensed for use in CAB. An overall analysis of these studies showed that, although bicalutamide 50 mg/day had clinical benefits, it was inferior to castration in terms of overall survival (median difference 97 days) (50). Subsequent
dose-ranging studies established that bicalutamide 150 mg once daily achieved a PSA response similar to that seen with castration while maintaining a good tolerability profile (51). Accordingly, the 150 mg dosage was chosen for further evaluation as both primary and adjuvant monotherapy.

As primary monotherapy, bicalutamide 150 mg/day has been compared to medical or surgical castration in two large prospective randomized trials with identical study design, including a total of 1,435 patients with locally advanced M0 or M1 CaP (51). A pooled analysis showed:

- In M1 patients, an improvement in overall survival with castration, although the difference in median survival between the groups was only 6 weeks (52); a further post-hoc analysis showed a survival benefit only for patients with higher PSA level (> 400 ng/mL) at study entry (53).
- In M0 patients, no significant difference was noted in overall survival (54,55).

In two smaller randomized trials, high-dose bicalutamide was compared to CAB. In the first trial (251 patients with predominantly M1 stage), no difference in overall survival was apparent (56). In the second trial (220 patients with M0 and M1 stage), there was no difference in overall survival for well- or moderately well-differentiated tumours (57) (level of evidence: 1b), but both studies were underpowered.

As for the adjuvant setting, the ongoing Early Prostate Cancer Programme (a study comprising 3 different clinical trials of similar design and including 8,113 patients worldwide) was designated to evaluate the efficacy and tolerability of high-dose (150 mg/day) bicalutamide versus placebo given in addition to standard primary care (i.e. radical prostatectomy, radiotherapy and “watchful waiting”) in localized or locally advanced CaP. The first combined analysis of the programme showed that, after a median follow-up of 3 years, adjuvant bicalutamide provided a reduction of 42% in the risk of objective disease progression compared to standard care alone (58). After a median follow-up of 5.4 years, it was shown that the positive effects of bicalutamide were obvious in patients with locally advanced disease (stage M0), whereas for patients with localized disease survival appeared to be reduced as compared to those receiving placebo (59).

In conclusion, high-dose bicalutamide has emerged as an alternative to castration for patients with locally advanced (M0) and in highly selected, well-informed cases of M1 CaP, but should be avoided in patients with localized CaP.

11.3.3  Combination therapies

11.3.3.1 Complete androgen blockade (CAB)

Although serum testosterone levels are reduced by up to 95% by castration, the intraprostatic androgen stimulus is sustained by the conversion of the circulating androgens of adrenal origin into DHT within the prostate cells. The action of these adrenal androgens is blocked by the addition of an antiandrogen to either surgical or pharmacological castration, a concept known as complete (or maximal or total) androgen blockade (CAB).

A plethora of studies evaluating CAB over monotherapy have been carried out with contrasting results. From the most recent systematic reviews and meta-analyses it appears that at a follow-up of 5 years, CAB provides a small survival advantage (less than 5%) when compared to monotherapy (60-64, level of evidence: 1a), even if some the largest trials included are methodologically flawed (65). It remains debatable whether this small advantage, if any, can be meaningful when applied to everyday clinical practice.

11.3.3.2 Minimal androgen blockade (or peripheral androgen blockade)

This derives from the combination of finasteride and a non-steroidal antiandrogen. The rationale behind the combination is that finasteride reduces intraprostatic levels of DHT by inhibiting 5α-reductase while antiandrogen competes with the binding of the residual DHT to its receptor. The result is that testosterone levels are maintained within normal ranges to ensure an acceptable sexual function and a reasonable quality of life (QoL).

In several phase II trials (66-70), the association of finasteride and flutamide, either in a concomitant or sequential regimen, has been evaluated in terms of PSA-response rate in patients with advanced or biochemically recurrent CaP. Notwithstanding the small sample and short follow-up, the overwhelming majority of patients experienced a substantial decline in PSA (by up to 96% compared to the level at entry). An update of one of these studies, at a long-term follow-up, reported on stronger endpoints, such as castration-free survival (median: 37 months), androgen-independent CaP-free survival (median: 48.6 months) and overall survival rate (65% at 5 years); the conclusion was that combination therapy can induce an overall period of hormone-responsive disease exceeding 4 years (71). In all these trials, sexual function was reported to be preserved in the great majority (55% to 86%) of men.

The preliminary data make this treatment option most attractive in the management of patients for whom QoL is the primary issue. However, while awaiting the results of follow-up and larger controlled trials, the treatment is still regarded as investigational.

11.3.3.3 Intermittent versus continuous androgen deprivation therapy (ADT)

For reasons that as yet remain unclear, long-term CAB which stimulates prostate cell apoptosis fails to eliminate the entire malignant cell population, so that after a variable period (averaging 24 months) the tumour
inevitably relapses being characterized by an androgen-independent state of growth. Experimental data indicate that androgen-independent progression may begin early after the administration of hormonal therapy, coinciding with the cessation of androgen-induced differentiation of stem cells (72). It is therefore theoretically possible that if androgen deprivation is stopped prior to the progression of androgen-independent cells, any subsequent tumour growth would then be solely sustained by the proliferation of androgen-dependent stem cells which should be susceptible once again to androgen withdrawal; in this way, cyclic ADT would delay the emergence of the androgen-independent clone. Thus, intermittent ADT may result in two other benefits, namely the preservation of QoL in the off-therapy periods and the reduction of cost.

Several phase II trials have demonstrated the feasibility of intermittent androgen blockade (IAB) in metastatic or biochemically recurrent disease, with PSA-response rates and symptom improvement similar to that of CAB, but phase III prospective, randomized controlled trials are still underway and data on survival endpoints and QoL are not mature (73).

In conclusion, although IAB is at present widely offered to patients with CaP in various clinical settings, its status should be regarded as investigational.

11.3.3.4 Immediate versus deferred ADT

The most appropriate time to introduce hormonal therapy in patients with advanced CaP is still controversial, in particular, whether ADT for locally advanced and asymptomatic metastatic disease delivered immediately at diagnosis favourably influences survival and QoL compared to ADT deferred while signs and symptoms of clinical progression remain a matter of debate.

The dispute derives from the lack of properly conducted, randomized, controlled trials, with many of them being methodologically flawed due to small size and underpowering and heterogeneity of patients enrolled as having advanced CaP (locally advanced, nodal and metastatic stage of disease), as well as variability in the hormone treatments administered and of follow-up schedules and modalities used.

Bearing in mind these limitations, evidence on immediate versus deferred ADT is provided by three systematic reviews of the literature (one of which is a meta-analysis). The Agency for Health Care Policy and Research report indicated that a possible survival advantage for early ADT existed in single studies where hormone treatment was the primary therapy while the combined analysis showed no significant benefit. Furthermore, androgen suppression was shown to be most cost-effective if initiated after patients experienced symptoms from metastatic disease (60,74). The Cochrane Library review extracted four good-quality randomized controlled trials (namely, VACURG I & II studies (9,10), the MRC trial (75) and the Eastern Cooperative Oncology Group (ECOG) 7887 study (76)), which were all conducted in the pre-PSA era and included patients with advanced CaP who received early versus deferred ADT as primary therapy or adjuvant to radical prostatectomy, but not to radiotherapy. According to the analysis, early androgen suppression significantly reduces disease progression and complication rates due to the progression itself, but does not improve cancer-specific survival and provides a relatively small benefit in overall survival with an absolute risk reduction of 5.5%, which does not become evident until after 10 years (77). Based on a systematic review of the literature, the recently published American Society of Clinical Oncology guidelines on the initial hormonal treatment for androgen-sensitive metastatic, recurrent or progressive CaP concluded that no recommendation can be made as to when to start hormonal therapy in advanced asymptomatic CaP until data from studies using modern diagnostic and biochemical tests and standardized follow-up schedules become available (78).

For asymptomatic patients with locally or regionally advanced CaP who undergo radiotherapy, there is good evidence from several randomized controlled trials that concomitant and/or adjuvant hormonal therapy provides longer time-to-disease progression and/or longer overall survival than radiotherapy alone followed by androgen suppression at progression (level of evidence: 1b) (79-82).
11.4  Indications for hormonal therapy (Table 9)

Table 9: Indications for hormonal therapy

<table>
<thead>
<tr>
<th>Castration</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>• M1 symptomatic</td>
<td>To palliate symptoms and to reduce the risk for potentially catastrophic sequelae of advanced disease (spinal cord compression, pathological fractures, ureteral obstruction, extra-skeletal metastasis) (level of evidence: 3)</td>
</tr>
<tr>
<td>• M1 asymptomatic</td>
<td>Immediate castration to defer progression to a symptomatic stage and prevent serious disease progression-related complications (level of evidence: 1b)</td>
</tr>
<tr>
<td>• N+ Immediate castration</td>
<td>Refs. 76,79 (level of evidence: 1b)</td>
</tr>
<tr>
<td>• Locally advanced M0</td>
<td>Immediate castration to improve cancer-free survival (level of evidence: 1b)</td>
</tr>
<tr>
<td>• Locally advanced symptomatic</td>
<td>Ref. 84 (level of evidence: 4)</td>
</tr>
<tr>
<td>• Locally advanced asymptomatic</td>
<td>Ref. 85</td>
</tr>
</tbody>
</table>

**Antiandrogens**

| Short-term administration | To reduce the risk of the “flare up” phenomenon in patients with advanced metastatic disease who are to receive an LHRH agonist (level of evidence: 1b) |
| Non-steroidal antiandrogens | Primary monotherapy as an alternative to castration in patients with locally advanced CaP (level of evidence: 1b). |

11.5  Contraindications for various therapies (Table 10)

Table 10: Contraindications for various therapies

<table>
<thead>
<tr>
<th>Therapy</th>
<th>Contraindications</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Bilateral orchiectomy</td>
<td>Psychological reluctance to surgical castration</td>
</tr>
<tr>
<td>• Oestrogens</td>
<td>Known cardiovascular disease</td>
</tr>
<tr>
<td>• LHRH agonists</td>
<td>Patients with metastatic disease at high risk for clinical “flare up” phenomenon</td>
</tr>
<tr>
<td>• Antiandrogens</td>
<td>Localized CaP as primary therapy</td>
</tr>
<tr>
<td></td>
<td>Known hepatic dysfunction</td>
</tr>
</tbody>
</table>

11.6  Outcome

Outcome depends on the stage and grade of disease at diagnosis. In M1 cases, the median overall survival ranges between 28 and 53 months (60); only 7% of patients with metastatic cancer treated with hormonal therapy are reported to live 10 years or more (90). Survival is likely to depend on PSA level at diagnosis, Gleason score, volume of metastatic disease and presence of bony symptoms. In locally advanced M0 patients, the median overall survival is frequently reported exceeding 10 years (61).

11.7  Side-effects, QoL and cost of hormonal therapy

Many patients with CaP for whom long-term ADT is indicated are still young and physically and sexually active, so QoL is an issue of paramount importance when considering the various hormonal treatment options. In view of this, in selected patients, monotherapy with a non-steroidal antiandrogen (i.e., bicalutamide) is gaining increasing interest due to its appeal in maintaining normal (or even higher) serum testosterone levels and in showing a good tolerability profile.

11.7.1  Side-effects

The number of deleterious side-effects during long-term androgen deprivation therapy has been well known for years (Table 11). Some of these can have a detrimental effect on QoL, especially in young men, while others may contribute to increased risk for serious health concerns associated with age.
### Table 11: Side-effects of hormonal treatment (adapted from Higano, et al. [91])

<table>
<thead>
<tr>
<th>Side-effects of therapy</th>
<th>Treatment/prophylaxis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Castration</strong></td>
<td></td>
</tr>
<tr>
<td>• Loss of libido</td>
<td>None</td>
</tr>
<tr>
<td>• Erectile dysfunction</td>
<td>Phosphodiesterase-5 (PDE5)-inhibitors, intracavernosal injection (ICI), vacuum device</td>
</tr>
<tr>
<td>• “Hot flashes” (55-80% of patients during androgen deprivation therapy)</td>
<td>Diethylstilboestrol (DES), cyproterone acetate (CPA), venlafaxine, clonidine</td>
</tr>
<tr>
<td>• Gynaecomastia and breast pain (49-80% DES, 50% complete androgen blockade (CAB), 10-20% castration)</td>
<td>Prophylactic radiotherapy, mammectomy, tamoxifen, aromatase inhibitors</td>
</tr>
<tr>
<td>• Increase in body fat</td>
<td>Exercise</td>
</tr>
<tr>
<td>• Muscle wasting</td>
<td>Exercise</td>
</tr>
<tr>
<td>• Anaemia (severe in 13% CAB)</td>
<td>Erythropoietin (EPO)</td>
</tr>
<tr>
<td>• Decrease in bone mineral density (not DES)</td>
<td>Exercise, calcium+ vitamin D, bisphosphonates</td>
</tr>
<tr>
<td>• Cognitive decline (not DES)</td>
<td>None</td>
</tr>
<tr>
<td><strong>Oestrogens</strong></td>
<td></td>
</tr>
<tr>
<td>• Cardiovascular toxicity (acute myocardial infarction, congestive heart failure, cerebrovascular accident, deep vein thrombosis, pulmonary embolism)</td>
<td>Parenteral administration, anticoagulants</td>
</tr>
<tr>
<td><strong>Antiandrogens</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Steroidal</strong></td>
<td></td>
</tr>
<tr>
<td>• Pharmacological side-effects: loss of libido, erectile dysfunction, rarely gynaecomastia</td>
<td></td>
</tr>
<tr>
<td>• Non-pharmacological side-effects: see individual drugs</td>
<td></td>
</tr>
<tr>
<td><strong>Non-steroidal</strong></td>
<td></td>
</tr>
<tr>
<td>• Pharmacological side-effects: gynaecomastia(49-66%), breast pain (40-72%), ‘hot flashes’(9-13%)</td>
<td>Prophylactic radiotherapy, mammectomy, tamoxifen, aromatase inhibitors</td>
</tr>
<tr>
<td>• Non-pharmacological side-effects: see individual drugs</td>
<td></td>
</tr>
</tbody>
</table>

11.7.2 Quality of life (QoL)

Data on QoL during hormone treatment are scant because of lack of solid evidence. The only large prospective randomized study is a double-blind placebo-controlled trial including 739 patients with M1 CaP comparing orchietomy plus flutamide versus orchietomy plus placebo. QoL was assessed in the first 6 months of treatment and the combined therapy resulted in lower QoL, with statistically significant differences in two QoL parameters, namely more frequent diarrhoea and worse emotional functioning than castration alone (92).

A prospective non-randomized observational study, including 144 patients with locally advanced CaP or PSA failure after definitive local treatment, showed that patients who received immediate ADT (by means of bilateral orchietomy, LHRH agonist or CAB) reported a lower overall QoL (increased fatigue, emotional distress and decreased physical functioning) than patients in the deferred hormone treatment arm (93, level of evidence: 2a).

A retrospective non-randomized study, including 431 patients with stage CaP who received orchietomy or LHRH agonists as their primary therapy within 12 months after initial diagnosis, assessed health-related quality of life (HRQoL) outcomes at 12 months’ follow-up. Men receiving LHRH agonists reported more worry and physical discomfort and a poorer overall health and were less likely to believe themselves free of cancer compared with orchietomized patients; the stage at diagnosis had no significant independent effect on health outcome. However, the study was insufficiently powered (94, level of evidence: 2b).

A recent, small, randomized controlled trial evaluated HRQoL of patients with non-localized CaP allocated to leuprolelin, goserelin, CPA and no treatment at 1-year follow-up; both sexual and cognitive function significantly declined in men on all forms of androgen suppression, while emotional distress significantly increased in those assigned to CPA and no treatment (95, level of evidence: 1b).

As for antiandrogen monotherapy, QoL was evaluated in the previously mentioned combined studies of bicalutamide monotherapy by means of a validated questionnaire covering 10 domains (sexual interest, sexual function, physical capacity, emotional well-being, vitality, social function, activity limitation, pain, bed disability and overall health). Separate analyzes of data for M0 and M1 patients were performed at 12-month
follow-up, and in both patient categories bicalutamide showed a significant advantage over castration in the domains of physical capacity and sexual interest (55, level of evidence: 1b). A further post-hoc analysis, including only the patients with sexual interest at study entry, showed that significantly more patients receiving bicalutamide 150 mg/day maintained their interest in sex and felt that they were still sexually attractive compared with those randomized to castration (96). Data on QoL are also available from the early report of the study of Boccardo, et al. (97) and support the findings of the two larger combined trials, in that more men in the bicalutamide group than in the castration group reported a preserved libido and erectile function. Furthermore, a recent, small, prospective randomized trial, including 103 patients with localized or locally advanced CaP who received bicalutamide 150 mg/day or medical castration, evaluated the changes in bone mineral density after 96 weeks of treatment and showed that bone mineral density is maintained with bicalutamide therapy (89, level of evidence: 1b). The most common side-effects during non-steroidal antiandrogen monotherapy are gynaecomastia and breast pain, which are caused by an imbalance in the androgen/oestrogen ratio within the breast tissue; in the bicalutamide studies, these events were reported by up to 66% and 73% of patients, respectively, but they were generally well tolerated, with a low withdrawal rate from therapy (58).

11.7.3 Cost-effectiveness of hormonal therapy options
A recent formal meta-analysis and literature review evaluated the cost-effectiveness of various long-term androgen suppression options in advanced CaP (i.e., bilateral orchiectomy, DES, LHRH-agonist, non-steroidal antiandrogen monotherapy, CAB with a non-steroidal antiandrogen). For the analysis, a sophisticated statistical model was generated, assuming the base case at entry to be a 65-year-old man with a clinically evident, local recurrence of CaP and no distant metastases, followed for a 20-year time horizon. The study concluded that, for men who can accept it, bilateral orchiectomy is the most cost-effective form of ADT providing a higher quality-adjusted survival, while CAB is the least economically attractive option, yielding small health benefits at high relative cost. Furthermore, the greatest QoL gains and least costs may be obtained by starting ADT when symptoms from distant metastases have occurred (74, level of evidence: 1a).

11.8 SUMMARY OF HORMONAL THERAPY
1. In advanced CaP, ADT delays progression, prevents potentially catastrophic complications and effectively palliates symptoms, but does not prolong survival (level of evidence: 1b)
2. In advanced CaP, all forms of castration as monotherapy (orchiectomy, LHRH and DES) have equivalent therapeutic efficacy (level of evidence: 1b)
3. Non-steroidal antiandrogen monotherapy (e.g. bicalutamide) is an effective alternative to castration in patients with locally advanced disease (level of evidence: 1b)
4. In advanced CaP, the addition of a non-steroidal antiandrogen to castration (CAB) results in a small advantage in overall survival over castration alone but is associated with increased adverse events, reduced QoL and high costs (level of evidence: 1a)
5. Intermittent and “minimal” ADT should still be regarded as experimental therapies (level of evidence: 3)
6. In advanced CaP, immediate (given at diagnosis) androgen suppression significantly reduces disease progression and complication rate due to progression itself compared to deferred (delivered at symptomatic progression) androgen deprivation (level of evidence: 1b)
7. Bilateral orchiectomy may be the most cost-effective form of ADT, especially if initiated after occurrence of symptoms from metastatic disease (level of evidence: 3).

11.9 REFERENCES


48. Pavone Macaluso M. Flutamide monotherapy versus combined androgen blockade in advanced prostate cancer. Interim report of an Italian multicentre, randomized study. SIU 23rd Congress 1994;354A.


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<table>
<thead>
<tr>
<th>Stage</th>
<th>Treatment</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1a</td>
<td>Watchful waiting</td>
<td>Standard treatment for well- and moderately, differentiated tumours and &lt; 10-year life expectancy. In patients with &gt; 10-year life expectancy, re-staging with TRUS and biopsy is advised (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radical prostatectomy</td>
<td>Optional in young patients with a long life expectancy, especially for poorly differentiated tumours (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy</td>
<td>Optional in younger patients with a long life expectancy, especially for poorly differentiated tumours. Higher complication risks after TURP, especially with interstitial radiation (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Hormonal</td>
<td>Not an option (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>Not an option (grade C recommendation)</td>
</tr>
<tr>
<td>T1b-T2b</td>
<td>Watchful waiting</td>
<td>Asymptomatic patients with well- and moderately, differentiated tumours and a life expectancy &lt; 10 years. Patients who do not accept treatment-related complications (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radical prostatectomy</td>
<td>Standard treatment for patients with life expectancy &gt; 10 years who accept treatment-related complications (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy</td>
<td>Patients with a life expectancy &gt; 10 years who accept treatment-related complications. Patients with contraindications for surgery. Unfit patients with 5-10 years of life expectancy and poorly differentiated tumours (combination therapy is recommended; see below) (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Hormonal</td>
<td>Symptomatic patients who needs palliation of symptoms unfit for curative treatment. (grade C recommendation). Antiiandrogens are associated with poorer outcome in comparison with watchful waiting and are not recommended (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>Neoadjuvant hormonal therapy (NHT) + radical prostatectomy: no proven benefit (grade A recommendation) NHT + radiotherapy: better local control. No proven survival benefit (grade B recommendation). Hormonal (3 years) + radiotherapy: better than radiotherapy in poorly differentiated tumours (grade A recommendation)</td>
</tr>
<tr>
<td>T3-T4</td>
<td>Watchful waiting</td>
<td>Option in asymptomatic patients with T3, well-differentiated and moderately differentiated tumours, and a life expectancy &lt; 10 years (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radical prostatectomy</td>
<td>Optional for selected patients with T3a and a life expectancy &gt; 10 years (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy</td>
<td>T3 with &gt; 5-10 years of life expectancy. Dose escalation &gt; 70 Gy seems to be of benefit. If this is not available, a combination with hormonal therapy could be recommended (see below) (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Hormonal</td>
<td>Symptomatic patients, extensive T3-T4, high PSA level (&gt; 25 ng/mL), unfit patients. Better than watchful waiting (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>Radiotherapy + hormonal seems better than radiotherapy alone (grade A recommendation). NHT + radical prostatectomy: no proven benefit (grade B recommendation)</td>
</tr>
<tr>
<td>N+, M0</td>
<td>Watchful waiting</td>
<td>Asymptomatic patients. Patient driven. May have worse survival (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radical prostatectomy</td>
<td>No standard option (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy</td>
<td>No standard option (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Hormonal</td>
<td>Standard therapy (grade A recommendation)</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>No standard option. Patient driven (grade B recommendation)</td>
</tr>
<tr>
<td>M+</td>
<td>Watchful waiting</td>
<td>No standard option. May have worse survival/more complications than with immediate hormonal therapy (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radical prostatectomy</td>
<td>Not an option (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Radiotherapy</td>
<td>Not an option (given for cure) (grade C recommendation)</td>
</tr>
<tr>
<td></td>
<td>Hormonal</td>
<td>Standard therapy. Symptomatic patients should not be denied treatment (grade B recommendation)</td>
</tr>
<tr>
<td></td>
<td>Combination</td>
<td>Not an option (grade C recommendation)</td>
</tr>
</tbody>
</table>
13 FOLLOW-UP: AFTER TREATMENT WITH CURATIVE INTENT

13.1 Definition
Curative treatment is defined as radical prostatectomy or radiotherapy, either by external beam radiation or an interstitial technique, or any combination of these. Alternative treatment options that are not fully established, such as CSAP, HIFU or RITA, are outside the scope of this chapter.

13.2 Why follow-up?
The first question to be answered is: ‘If failure after curative treatment is so common, are follow-up efforts worthwhile?’ The number of patients who will have a detectable PSA level after radical prostatectomy varies between published series. The Johns Hopkins group reported 16%, 26% and 34% biochemical progression after 5, 10 and 15 years, respectively. At the Cleveland Clinic, the 5-year biochemical progression rate was as high as 39% in stage T1-T2 CaP (1,2). Similar data have been presented by European centres (3). It has also been shown that the risk of relapse after radical prostatectomy can persist even after 5 years, suggesting that follow-up should be continued for a longer time period (1,4). Following radiotherapy, there is a similar course of events. A considerable proportion of patients will have a rising PSA level, and disease recurrences will continue to become obvious, even after 15 years of follow-up (5-7).

The answer to the first question is therefore definitely ‘Yes’; recurrences will occur in a substantial number of patients who received treatment with intent to cure.

The second question to be answered is: ‘What is the reason for follow-up?’ Reasons may vary depending on the treatment given, patient age, co-morbidity and the patient’s own will. In general, patients who receive curative therapy may be followed-up for any of the following reasons:

• Good responsible patient care
• Possibility of second-line treatment with curative intent
• Possibility of early hormonal therapy after failure
• As part of a study protocol.

Chapter 15 discusses treatment options.

13.3 How to follow-up?
The procedures indicated at follow-up visits vary depending on the clinical situation. The examinations discussed below are routinely used for the detection of CaP progression or residual disease. The PSA level, and eventually DRE, are the only tests that need to be carried out routinely. A disease-specific history should be mandatory at every follow-up visit and should include psychological aspects, signs of disease progression and treatment-related complications. The examinations used for the evaluation of treatment-related complications must be individualized and are beyond the scope of these guidelines. The examinations used most often for cancer-related follow-up after curative surgery or radiation treatment are discussed below.

13.3.1 PSA monitoring
The measurement of PSA level is a cornerstone of follow-up after curative treatment. There is a difference in what can be expected after radical prostatectomy and radiotherapy, but PSA recurrence nearly always precedes clinical recurrence after either treatment, in some cases by many years (1,5,8-10). It is recommended that the finding of a single, elevated, serum PSA level should be re-confirmed before treatment is altered.

It is worth pointing out that the use of hormonal therapy before, during, or after curative treatment may make PSA unreliable as a tumour marker for follow-up. It has been shown that a 3-month treatment course with LHRH analogue (LHRHa) prior to radical prostatectomy can delay PSA progression by approximately 1 year, without obvious impact on later progression-free survival (11). A 3-year course of LHRHa, as advocated for bulky localized CaP treated with radiotherapy, may well have an even larger influence on PSA level as a follow-up tool (12,13).

13.3.2 Definition of PSA progression
The level of PSA at which to define treatment failure differs between radical prostatectomy cases and radiation-treated cases. Following radical retropubic prostatectomy, two consecutive values of 0.2 ng/mL or greater appear to represent an international consensus defining recurrent cancer (14,15). Following radiation therapy, a reasonable definition of biochemical relapse is three consecutive increases according to the recommendation of ASTRO (16).
13.3.3 PSA monitoring after radical prostatectomy

PSA is expected to be undetectable within 3 weeks after a successful radical prostatectomy (17). A persistently elevated PSA level means that PSA-producing tissue remains in the body. In patients treated with radical prostatectomy, this is generally thought to be residual cancer due to either micrometastases that were not detected or undetectable beforehand, or residual disease in the pelvis possibly due to positive surgical margins.

A rapidly increasing PSA level (high PSA velocity, short PSA doubling time) indicates rather distant metastases, while a later and slowly increasing concentration of PSA is most likely to indicate local disease recurrence. The time to PSA recurrence and tumour differentiation are also important predictive factors distinguishing between local and systemic recurrence (18,19). Both local treatment failure and distant metastases have been shown to occur with undetectable PSA levels. This is very rare and occurs almost only in patients with unfavourable pathology (undifferentiated tumours) (20,21).

This means that, in patients with a relatively favourable pathology (< pT3, pN0, Gleason score < 8), PSA measurement, together with the disease-specific history, could stand as the single test in follow-up after radical prostatectomy. The PSA cut-off point recommended should be no lower than 0.2 ng/mL. It has been shown that patients with a PSA level between 0.1 ng/mL and 0.2 ng/mL after radical prostatectomy had neither clinical nor biochemical disease progression (22). Furthermore, no adjuvant treatment given at an even earlier stage has proved to be beneficial to patients with PSA relapse. Therefore, the use of an ultra-sensitive PSA assay is not justified for routine follow-up after radical prostatectomy. If ongoing randomized trials show that early adjuvant treatment after radical prostatectomy improves survival, this issue should be reconsidered.

13.3.4 PSA monitoring after radiation therapy

The PSA level falls slowly after radiotherapy compared with radical prostatectomy. The optimal cut-off value for a favourable PSA nadir after radiotherapy is somewhat controversial. Achieving a PSA nadir of less than 1 ng/mL seems to be associated with a favourable outcome, at least within the 3-5 year perspective (23). Lately, however, it has been suggested that this nadir level should be reduced to less than 0.5 ng/mL. This is because only 4% of treated patients with a nadir of less than 0.5 ng/mL failed therapy after 40 months of follow-up compared with 26% of those with a nadir of 0.6-1.0 ng/mL (24). The interval before reaching the nadir PSA may be very long and can sometimes take up to 3 years or more. Also, after radiotherapy, the PSA doubling time has been shown to correlate to the site of recurrence; patients with local recurrence had a doubling time of 13 months compared to 3 months for those with distant failure (7).

There is a consensus that a rising PSA level is an early sign of treatment failure. This has led ASTRO to define failure after radiation therapy as three consecutive rises in PSA level, irrespective of the nadir value (16). It is important to realize that this definition may cause outcomes that are difficult to interpret in certain situations. Depending on the timing of the PSA measurements, the time before a biochemical progression is ultimately recognized may be delayed by several years. Also, these criteria are hard to use after adjuvant/neoadjuvant hormonal therapy, commonly used together with radiotherapy.

13.3.5 Digital rectal examination (DRE)

DRE is performed to assess whether or not there is any sign of local disease recurrence. It is very difficult to interpret the findings of DRE after curative therapy, especially after radiotherapy. A newly detected nodule should raise the suspicion of local disease recurrence.

As mentioned previously, a local disease recurrence after curative treatment is possible without a concomitant rise in PSA level (20,21). However, this has only been proven in patients with unfavourable pathology, i.e. those with undifferentiated tumours. Thus, PSA measurement and DRE comprise the most useful combination of tests as first-line examination in follow-up after radiotherapy or radical prostatectomy, but PSA measurement may well be the only test in cases with favourable pathology.

13.3.6 Transrectal ultrasonography (TRUS) and biopsy

TRUS and biopsy have no place in the routine follow-up of asymptomatic patients and nowadays only rarely after biochemical failure. TRUS cannot stand alone as a diagnostic tool, but must usually be combined with biopsy to establish the presence of local disease recurrence. The purpose of the investigation is to confirm a histological diagnosis of local disease recurrence. It is only warranted if the finding of a local recurrence affects the treatment decision (see Section 15 for a more detailed discussion).

13.3.7 Bone scintigraphy

The purpose of bone scintigraphy is to detect skeletal metastases. It is not recommended for the routine follow-up of asymptomatic patients, but may be indicated in individuals with elevated PSA levels for whom the findings will affect the treatment decision. It is also indicated in patients with symptoms arising from the skeleton, since metastatic disease may occur even if PSA is undetectable (20,21).
13.3.8 Computed tomography (CT) or magnetic resonance imaging (MRI)

CT or MRI have no place in the routine follow-up of asymptomatic patients. They may be used selectively in the evaluation after biochemical failure before treatment decisions are made (see Section 15).

13.4 When to follow-up?

Most patients who fail treatment for CaP do so early, even if failure only becomes clinically obvious after years (1-7). The patient should therefore be followed-up more closely during the first years after treatment when the risk of failure is highest. PSA measurement, disease-specific history and DRE are recommended at the following intervals: 3, 6 and 12 months postoperatively, every 6 months thereafter until 3 years, and then annually.

The purpose of the first clinic visit is mainly to detect treatment-related complications and to assist patients in coping with the new situation. Tumour or patient characteristics may allow alterations to this schedule. For example, patients with poorly differentiated and locally advanced tumours or with positive margins may be followed-up more closely than those with a well-differentiated, intracapsular or specimen-confined tumour. Obviously, advanced age or associated co-morbidity may make further follow-up in asymptomatic patients superfluous.

13.5 GUIDELINES FOR FOLLOW-UP AFTER TREATMENT WITH CURATIVE INTENT

1. In asymptomatic patients, a disease-specific history and a serum PSA measurement supplemented by DRE are the recommended tests for routine follow-up. These should be performed at 3, 6 and 12 months after treatment, then every 6 months until 3 years, and then annually (grade B recommendation)

2. After radical prostatectomy, a serum PSA level of more than 0.2 ng/mL can be associated with residual or recurrent disease (grade B recommendation)

3. After radiation therapy, a rising PSA level, rather than a specific threshold value, is the most reliable sign of persistent or recurrent disease (grade B recommendation)

4. Both a palpable nodule and a rising serum PSA level can be signs of local disease recurrence (grade B recommendation)

5. Detection of local recurrence by TRUS and biopsy is only recommended if it will affect the treatment plan. In most cases TRUS and biopsy are not necessary before second-line therapy (grade B recommendation)

6. Metastasis may be detected by pelvic CT/MRI or bone scan. In asymptomatic patients, these examinations may be omitted if the serum PSA level is less than 30 ng/mL but data on this topic are sparse (grade C recommendation)

7. Routine bone scans and other imaging studies are not recommended in asymptomatic patients. If a patient has bone pain, a bone scan should be considered irrespective of the serum PSA level (grade B recommendation).

13.6 REFERENCES


14  FOLLOW-UP: AFTER HORMONAL TREATMENT

14.1  Introduction
A large proportion of patients treated with hormonal therapy have either metastatic or locally advanced tumours at diagnosis. This will affect the scheme of follow-up because biochemical failure is often associated with rapid symptomatic progression.

14.2  Why follow-up?
The main objectives of follow-up in these patients are:
  • To monitor the response to treatment
  • To ensure compliance with treatment
  • To detect potential complications of endocrine therapy
  • To guide the modalities of palliative symptomatic treatment at the time of hormonal escape.

However, the usefulness of complementary investigations at various stages of the disease must be clarified in order to avoid useless examinations and an excessive economic cost to the community. On the other hand, strict recommendations for follow-up procedures are only useful if effective therapeutic strategies can be offered to the patient in case of disease progression. To date, the issue of early versus late initiation of non-hormonal treatment in HRPC has still not been resolved so that follow-up should be performed on an individual
basis. Based on current knowledge, strict guidelines for follow-up procedures following hormonal therapy cannot be formulated.

14.3 How to follow-up?

14.3.1 PSA monitoring
Prostate-specific antigen is a good marker with which to follow the course of metastatic CaP and is more reliable than PAP. Many authors have studied the prognostic value of PSA (prediction of the duration of response to endocrine treatment) based on either the initial pre-treatment value or the PSA decrease during the first 3-6 months (1,2). The initial PSA level can reflect the extent of metastatic disease, although some poorly differentiated tumours do not secrete PSA. The prognostic value of the pre-treatment PSA value is variably assessed in the literature and should not be used to predict the duration of response to treatment (3).

Treatment response may be assessed utilizing the change in serum PSA level as a surrogate endpoint after hormonal treatment has been initiated. The PSA decrease can be evaluated in terms of the absolute PSA level at 3 months or 6 months, the nadir PSA during treatment, or the rate at which PSA decreases (2,4,5). The PSA value after 3 and/or 6 months of hormonal treatment has been reported as being related to prognosis (3,5-7). However, this criterion has no absolute value in any individual patient (5,8). The subgroup of patients with a normal or undetectable PSA level at 3 and 6 months corresponds to the group with the highest probability of long-lasting response to endocrine treatment.

After the initial phase of response to endocrine treatment, patients should be regularly monitored in order to detect and treat any complications of endocrine escape, as clinical disease progression occurs after a median interval of about 12-18 months of treatment in patients with stage M1 disease. It is well established that regular PSA control in asymptomatic patients allows the earlier detection of biochemical escape, as the rise in PSA level usually precedes the onset of clinical symptoms by several months (1,9,10). However, it must be stressed that PSA level is not a reliable marker of escape and cannot stand alone as a follow-up test. Clinical disease progression with normal PSA levels has been reported to occur in 15-34% of cases (9,11). Two mechanisms could explain the occurrence of tumour progression despite a normal PSA level in the context of androgen suppression. Firstly, antiandrogen activity and the fall in PSA level during endocrine treatment are not always proportional to the reduction in tumour volume (10,12-16). Secondly, the proportion of poorly differentiated cells in the tumour, which secrete less PSA, increases during endocrine treatment (17-20).

14.3.2 Creatinine, haemoglobin and liver function monitoring
Creatinine monitoring has some value because it can detect upper urinary tract obstruction in cases of advanced cancer that might need to be relieved by, for example, percutaneous nephrostomy or double J-stent. Haemoglobin and liver function tests could suggest disease progression and/or toxicity of hormonal treatment, which can lead to interruption of hormonal treatment (i.e. liver toxicity from non-steroidal antiandrogens).

The fact that haemoglobin levels will decrease by about 20% with androgen deprivation has to be taken into consideration (21).

Alkaline phosphatase and its bone-specific isoenzymes may be used to monitor patients with stage M1b disease. These markers have the advantage of not being directly influenced by hormonal therapy compared with PSA. It should be remembered that increases in serum concentrations of alkaline phosphatase might also be due to osteoporosis induced by androgen deprivation (22). In this scenario, the determination of bone-specific alkaline phosphatase might be helpful.

14.3.3 Prostatic acid phosphatase (PAP) monitoring, bone scan, ultrasound and chest X-ray
The monitoring of PAP levels no longer has any value since the introduction of PSA measurement (9). In routine practice, asymptomatic patients with a normal PSA level should not have a bone scan at regular intervals as disease progression is more reliably detected by PSA monitoring, which also has a lower cost (23-25).

Moreover, the interpretation of bone scans is sometimes difficult, and the appearance of a new site of uptake or deterioration of pre-existing lesions in an asymptomatic patient does not modify the therapeutic approach.

In cases where there is a clinical or laboratory suspicion of disease progression, a chest X-ray or renal or hepatic ultrasound may be indicated as well as TRUS. However, these examinations are not recommended for routine use in asymptomatic patients. In hormone-refractory disease, follow-up examinations should be individualized with the aim of maintaining the patient's quality of life.

14.4 When to follow-up?
After initiation of hormonal treatment, it is recommended that patients be followed-up at 3 and 6 months.
14.4.1 Stage M0 patients
If there is a good treatment response, i.e. symptomatic improvement, good psychological coping, good treatment compliance and a serum PSA level of less than 4 ng/mL, follow-up visits are scheduled every 6 months.

14.4.2 Stage M1 patients
If there is a good treatment response, i.e. good symptomatic improvement, good psychological coping, good treatment compliance and a serum PSA level of less than 4 ng/mL, follow-up is scheduled every 3-6 months. Patients on antiandrogen treatment may need closer follow-up as they might benefit from antiandrogen withdrawal at the time of disease progression.

14.4.3 Hormone-refractory patients
Patients whose disease progresses, or who do not respond according to the criteria mentioned above, warrant an individualized follow-up scheme.

14.5 GUIDELINES FOR FOLLOW-UP AFTER HORMONAL TREATMENT
1. Patients should be evaluated at 3 and 6 months after initiating treatment. Tests should include at least serum PSA measurement, DRE and careful evaluation of symptoms in order to assess the treatment response and the side-effects of treatments given (grade B recommendation)
2. Follow-up should be tailored for the individual patient, according to symptoms, prognostic factors and the treatment given (grade C recommendation)
3. In patients with stage M0 disease with a good treatment response, follow-up is scheduled every 6 months, and should include at least a disease-specific history, DRE and serum PSA determination (grade C recommendation)
4. In patients with stage M1 disease with a good treatment response, follow-up is scheduled for every 3-6 months. A minimal follow-up should include a disease-specific history, DRE and serum PSA determination, frequently supplemented with haemoglobin, serum creatinine and alkaline phosphatase measurements (grade C recommendation)
5. When disease progression occurs or if the patient does not respond to the treatment given, the follow-up needs to be individualized (grade C recommendation)
6. Routine imaging in stable patients is not recommended (grade B recommendation).

14.6 REFERENCES


15 TREATMENT OF BIOCHEMICAL FAILURE AFTER TREATMENT WITH CURATIVE INTENT

15.1 Background
Primary curative procedures, such as radical prostatectomy and radiotherapy, are well-established therapeutic options in the management of localized CaP. Technical advances in surgery and radiation therapy have improved therapeutic efficacy and decreased treatment-associated morbidity and toxicity, respectively. However, despite these improvements, there is still a significant risk of cancer recurrence after therapy and up to 27-53% of all patients undergoing radiation therapy or radical prostatectomy will develop local or distant recurrences within 10 years after initial therapy and 16-35% of patients will receive second-line treatment within 5 years of initial therapy (1-5,7).

15.2 Definitions
15.2.1 Definition of treatment failure
In previous years, treatment failure was defined as recurrence on DRE or the development of metastatic disease. Currently, treatment failure is defined as a rising PSA level based on a study of Pound et al. (6) who demonstrated that no patient followed for more than 5 years developed any recurrence without a concomitant PSA rise.

The level of PSA at which to define treatment failure differs between radical prostatectomy cases and radiation-treated cases. Following radical retropubic prostatectomy, two consecutive values of PSA ≥ 0.2 ng/mL appear to represent an international consensus defining recurrent cancer (7,8). However, the most appropriate definition of biochemical progression after radical prostatectomy is still uncertain. In a retrospective analysis of 2,782 men having undergone radical prostatectomy for clinically localized CaP, Amling et al. (9) determined the best PSA cut-off point to be used to define biochemical recurrence. The authors demonstrated that once PSA recurrence was detected, a subsequent increase in PSA was noted in 49%, 62% and 72% of patients who had PSA 0.2, 0.3, and 0.4 ng/mL, respectively. These data indicate that only half of patients with a PSA of 0.2 ng/mL will further progress and may initially be managed by surveillance. Following radiation therapy, a reasonable definition of biochemical relapse is three consecutive increases, according to the recommendation of ASTRO (10).
15.2.2 Definition of recurrence

- Following radical prostatectomy, two consecutive PSA values ≥ 0.2 ng/mL represent recurrent cancer.
- Following radiation therapy, three consecutive increasing PSA values above a previous nadir represent recurrent cancer.

15.3 Local or systemic relapse

With regard to further management, once PSA relapse has been diagnosed, it is of major importance to determine whether the recurrence has developed at local or distant sites. About 50% of the patients having undergone radical retropubic prostatectomy will have local disease and the remainder will have either distant disease alone, or distant and local disease (10).

Important parameters to help differentiate between local or distant relapse (Table 12) include: timing of PSA increase after surgery, PSA velocity, PSA doubling time (PSADT), pathohistological stage and Gleason score of the prostatectomy specimen. PSA elevations developing within the first 2 years following surgery are associated with distant recurrences (11). It was shown that a median PSADT of 4.3 months is associated with distant relapse, whereas a median PSADT of 11.7 months predicts local failure (12). According to a recent study (13), PSA velocity of < 0.75 ng/mL/year was observed in 94% of patients with local recurrence, whereas 56% of patients with distant metastases demonstrated a PSA velocity of > 0.75 ng/mL/year.

Table 12: Important clinical and pathohistological parameters predicting local and systemic relapse following radical prostatectomy

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Local recurrence</th>
<th>Systemic recurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interval to PSA relapse</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 1 year</td>
<td>7%</td>
<td>93%</td>
</tr>
<tr>
<td>1–2 years</td>
<td>10%</td>
<td>90%</td>
</tr>
<tr>
<td>&gt; 2 years</td>
<td>61%</td>
<td>39%</td>
</tr>
<tr>
<td>&gt; 3 years</td>
<td>74%</td>
<td>26%</td>
</tr>
<tr>
<td>PSA doubling time</td>
<td>11.7 months</td>
<td>4.3 months</td>
</tr>
<tr>
<td>Gleason Score</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2–4</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>5–6</td>
<td>55%</td>
<td>45%</td>
</tr>
<tr>
<td>7</td>
<td>39%</td>
<td>61%</td>
</tr>
<tr>
<td>8–10</td>
<td>11%</td>
<td>89%</td>
</tr>
<tr>
<td>Pathological stage</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organ confined (≤ pT2b)</td>
<td>40%</td>
<td>60%</td>
</tr>
<tr>
<td>pT3a, R0</td>
<td>54%</td>
<td>46%</td>
</tr>
<tr>
<td>pT3a, R1</td>
<td>48%</td>
<td>52%</td>
</tr>
<tr>
<td>pT3b</td>
<td>16%</td>
<td>84%</td>
</tr>
<tr>
<td>pTx N1</td>
<td>7%</td>
<td>93%</td>
</tr>
</tbody>
</table>

With radiation therapy, any continuously rising PSA following a nadir after radiation is an indicator for local recurrence, systemic metastatic spread or a combination of both (14-17). However, due to the well-known PSA bounce phenomenon, biochemical recurrence is defined by three consecutive PSA rises above the nadir level according to ASTRO guidelines. After radiotherapy, a late and slowly rising PSA is a sign of local failure only. Local recurrence is defined by a prostatic biopsy demonstrating malignant cells after 18 months or longer following initial radiotherapy, associated with a PSA rise, and no evidence of metastatic spread documented by CT/MRI and bone scintigraphy.

15.3.1 Definition of local and systemic failure

- Local failure following radical prostatectomy is predicted with an 80% probability by PSA increase > 3 years after radical prostatectomy, PSADT ≥ 11 months, Gleason score ≤ 6, stage ≤ pT3a pN0, pTx R1.
- Systemic failure following radical prostatectomy is predicted with > 80% accuracy by a PSA increase < 1 year after radical prostatectomy, PSADT of 4-6 months, Gleason score 8-10, stage pT3b, pTxN1.
- Local failure after radiation therapy is documented by a positive prostatic biopsy and negative imaging studies.
- Prostatic biopsy after radiation therapy is only necessary if local procedures such as salvage prostatectomy are indicated in an individual patient.
15.4 Evaluation of PSA progression

In recent years, most patients with PSA progression following initial therapy with curative intent underwent physical and sonographic examinations, radiographic studies or biopsies of the prostatic fossa and the vesicourethral anastomosis to confirm the recurrence identified by serological studies. For patients with asymptomatic PSA-only progression, the yield is very low and it has been shown by Lange et al. (13) that biochemical failure precedes clinical disease by 6-48 months.

In general, DRE is not useful in men with undetectable or very low PSA levels. In a recent study by Öbek et al. (18), it was shown that only 4 out of 72 patients (5.5%) with a PSA recurrence following radical prostatectomy had an abnormal DRE.

Traditionally, bone scans and abdominal CT scans have been applied to evaluate PSA elevations following primary treatment. Both imaging studies, however, are characterized by a low sensitivity and specificity and might be safely omitted in the routine work-up of relapsing patients. Recently, Cher et al. (19) studied 144 bone scans in 93 patients with PSA recurrence after radical retropubic prostatectomy. 122 patients had undergone radical prostatectomy without any hormonal treatment whereas 22 patients received either neoadjuvant or adjuvant ADT. Only 4.1% and 27% of the bone scintigrams were positive for metastatic disease; the lowest PSA associated with positive findings was 46 ng/mL in the absence of adjuvant androgen deprivation, whereas the lowest PSA value was 15.47 ng/mL in patients receiving hormonal therapy. The probability of a positive bone scan remains ≤ 5% until serum PSA reaches at least 40 ng/mL. Similar data have been achieved by other groups (20,21) demonstrating that patients with a true positive bone scan had an average PSA level of > 60 ng/mL and a PSA velocity of 22 ng/mL/year. On logistic regression analysis, PSA and PSA velocity predicted the findings on bone scan, and PSA velocity predicted the CT scan result. The probability of a positive bone scan and a positive CT scan was 9.4% and 14%, respectively, among the 132 patients with biochemical recurrence. However, there might be a slight difference between patients after radical retropubic prostatectomy compared to patients after radiation therapy, as demonstrated by Johnstone et al. (22) with 5% and 30% of the bone scans being positive, respectively.

In summary, bone scintigraphy and CT scans are of no additional diagnostic value unless the PSA serum levels are higher than 20 ng/mL or unless the PSA velocity is more than 20 ng/mL/year. Endorectal coil imaging has been described as a useful technique to detect local recurrences after radical prostatectomy (23). In a series of 48 patients, local recurrence was correctly identified in 81%, with the mean PSA at time of diagnosis being 2 ng/mL.

Positron emission tomography (PET) has been successfully applied in many human cancers for early identification of local or systemic recurrences. In CaP, there are only a few, but promising, data published on the clinical efficacy of PET in detecting local recurrences after radical prostatectomy (24). In a series of 31 patients with biochemical progression after radical prostatectomy, [11C]Acetate-PET demonstrated a high sensitivity and specificity for the detection of local recurrences if the PSA serum level was > 1 ng/mL.

Immunoscintigraphy using a radiolabelled monoclonal antibody based on prostate-specific membrane antigen for messenger RNA (PSMA) called 111 indium capromab penetide might represent an innovative diagnostic approach with an overall accuracy of up to 81% to detect the site of relapse in PSA-only recurrences following radical retropubic prostatectomy (25-28). Independent of the PSA serum concentration, the capromab pentetide scan shows a diagnostic yield of 60% to 80% and may help to stratify therapy according to the location of positive sites. In a recent study (28), 255 patients with PSA-only recurrence < 4.0 ng/mL after radical prostatectomy were investigated and capromab pentetide uptake was seen in 72% throughout the range of postoperative PSA serum levels (0.1-4.0 ng/mL). Approximately 31%, 42% and 25% of patients exhibited local uptake, locoregional and distant uptake, respectively, enabling a targeted therapy due to the differentiation of local versus systemic relapse.

It has been common practice to perform TRUS-guided biopsies of the prostatic fossa, the anastomosis or the prostate gland to exclude local recurrence after radical retropubic prostatectomy or radiation therapy. However, according to available studies, routine biopsy of the vesicourethral anastomosis appears not to be justified based on a verification rate of only 54% (29-33). Only in the presence of a palpable lesion or a hypoechoic lesion on transrectal ultrasound, can the diagnostic yield of the biopsy be improved to approximately 80%. Furthermore, there is a strong correlation between the positive biopsy rate and PSA serum concentrations (29-33). 28% and 70% of the biopsies were positive if the PSA level was below 0.5 ng/mL or greater than 2.0 ng/mL. It is common sense, nowadays, that routine anastomotic biopsy is not indicated and the use of PSA and PSADT is sufficient for clinical practice. In addition, PSA-free survival in biopsy-proven recurrences does not differ significantly as compared to PSA-only recurrences.

With regard to PSA relapses following radiation therapy, routine prostate biopsy should no longer be performed for the evaluation of PSA-only recurrences, according to an ASTRO consensus recommendation (34). However, prostate biopsy documenting local recurrence represents the main cornerstone in the decision making process for salvage radical prostatectomy in patients with rising PSA levels following a nadir after radiation therapy (35-37). It is a general recommendation to wait about 18 months and 3 months following radiation therapy or seeds and cryotherapy or HIFU, respectively.
15.5 **Diagnostic procedure in patients with PSA relapse**

1. Following radical prostatectomy, CT scans of the pelvis and abdomen are of low sensitivity and specificity in patients with PSA levels < 20 ng/mL or a PSA velocity of < 20 ng/mL/year.
2. Endorectal MRI or PET scans may help to detect local recurrences if PSA is > 1-2.0 ng/mL but is not yet part of routine clinical use.
3. If available, the capromab pendetide scan shows a diagnostic yield of 60% to 80% independent of the PSA level.
4. Following radiation therapy, local recurrence is documented by a positive biopsy ± 18 months after the procedure.

15.6 **Treatment of PSA-only recurrences**

The timing and mode of treatment of PSA-only recurrence after radical prostatectomy or radiation therapy remains controversial. After radical retropubic prostatectomy observation, radiation therapy to the prostatic bed, (complete) androgen blockade, intermittent androgen deprivation (IAD), combination of antiandrogens with 5α-reductase inhibitors and even early chemohormonal approaches are therapeutic options. The same therapeutic options may be applied for PSA recurrences following radiation therapy, in addition, salvage prostatectomy, cryotherapy and brachytherapy might be indicated in carefully selected patients.

15.6.1 **Radiation therapy for PSA-only recurrence after radical prostatectomy**

As confirmed by various studies the pre-radiation PSA level appears to be of critical importance in order to obtain optimal treatment results (38-46). Applying a pre-radiation cut-off of ≥ 2.5 ng/mL, Wu et al. (38) and Schild et al. (39) reported disease-free survival rates of 53% and 76% as compared to 8% and 26%, respectively, for patients with PSA levels ≥ 2.5 ng/mL. Forman et al. (40) demonstrated a disease-free survival rate of 83% vs 33% in patients with a PSA-only recurrence of less than 2.0 ng/mL and greater than 2.0 ng/mL, respectively. Nudell et al. (41) even reported a progression-free survival rate of 58% and 21% in patients having undergone radiation of the prostate bed if PSA serum levels were below 1.0 ng/mL or greater than 1.0 ng/mL, respectively. Based on these data, ASTRO has published a consensus paper recommending a dose of at least 64 Gy when the PSA level is < 1.5 ng/mL after radical retropubic prostatectomy (34).

These data of early salvage radiation therapy are corroborated by a recent paper (47) demonstrating a significant difference with regard to the 5-year biochemical-free and overall survival rates in patients being treated for PSA recurrence only or for palpable locally recurrent cancer. The 5-year biochemical-free and overall survival rates were 69% and 96% compared to 45% and 78%, respectively, in the group with palpable disease. However, data of prospective randomized trials are still lacking and all studies being performed lack long-term follow-up so that the impact on survival is unknown.

15.6.2 **Hormonal therapy**

In patients with a high pre-radical prostatectomy PSA > 20 ng/mL, a Gleason grade ≥ 7, an extensive positive surgical margin and extensive extraprostatic tumour growth (pT3b, pT3xN1) immediate hormonal therapy might be indicated (42-46). The impact of early ADT on long-term survival, however, is still unknown. In a retrospective observational multicentre study including 1,352 patients with PSA recurrence following radical prostatectomy (48), early ADT resulted in a significant reduction of the development of clinical metastases compared to delayed ADT. There was however no significant effect on long-term survival. These recommendations are corroborated by a recent study (50) demonstrating that none of the patients with a Gleason score 8, pT3b or pT3xN1 CaP remained disease-free following radiation therapy for PSA-only recurrence after radical prostatectomy.

It is difficult to make recommendations for the optimal therapeutic management for PSA-only recurrences following radical prostatectomy or radiation therapy because of the lack of prospective randomized trials. There are only very few studies analyzing the clinical utility of early androgen deprivation in locally advanced (M0) and metastatic PCA (50,51). It is believed that for the M0 category of the patients with pTnxN1 disease having undergone radical prostatectomy reflecting PSA-only recurrences, hormonal therapy would appear to be beneficial for some patients with a high probability of occult systemic metastases. There is some evidence that CAB has a pronounced survival benefit in patients with minimal metastatic disease so that patients with PSA-only recurrences might have a similar improved survival with combined androgen deprivation (52,53). Considering the speculative benefits, the side-effects of traditional hormonal therapy such as hot flushes, loss of libido, impotence, decreased muscular mass and osteoporosis must not be underestimated.

The use of antiandrogens alone might overcome these side-effects as demonstrated in recent studies. Although gynaecomastia and breast tenderness were the most predominant side-effects for the treatment of organ-confined and locally advanced CaP, the incidence of hot flushes, loss of libido and impotence was...
significantly lower than expected for LHRH-agonists and CAB (54). Furthermore, the risk of objective progression of the disease was significantly reduced in patients receiving bicalutamide 150mg (55). Antiandrogens may represent a viable alternative to other modes of androgen deprivation for the management of PSA-only recurrences especially in young and otherwise healthy men.

Non-traditional ways of using hormonal therapy for PSA-only recurrence include IAD and oral therapies combining antiandrogens with 5α-reductase inhibitors (56-63). In the setting of PSA-only recurrences, however, no prospective randomized trials and no clinical studies with sufficient data on long-term efficacy are available to justify a routine clinical application of IAD despite potential benefits. Summarizing the series in which PSA-only recurrences were treated by IAD (56-60), PSA threshold levels at study entry varied significantly as did the PSA level at discontinuation of hormonal therapy. Only the study by Tunn et al (60) involving 150 patients had an appropriate study design from which to draw important clinical conclusions. Patients were started on IAD for 9 months when the postprostatectomy PSA serum level was greater than 3.0 ng/mL and all patients reached a nadir of less than 0.5 ng/mL. IAD was restarted when PSA increased to more than 3.0 ng/mL; after a mean follow-up of 48 months and a mean duration of hormonal therapy of 26.6 months none of the patients had progressed to hormone-refractory disease.

In some studies, finasteride and flutamide have been combined to manage PSA-only recurrences since both agents work additively by blocking the intraprostatic conversion of testosterone to DHT and blocking the intracytoplasmic DHT receptor (61-63). In the latest report (63), including 73 patients, the application of finasteride (10 mg/day) and low-dose flutamide (250 mg/day) resulted in a mean PSA nadir of 1.35 ng/mL within 6 months. However, only 62% of the patients reached a PSA nadir of < 0.2 ng/mL. After a mean follow-up of 15 months, none of the patients had progressed to traditional hormonal therapy. However, longer follow-up of a larger patient cohort is needed and randomized phase III trials using modern antiandrogens with fewer gastrointestinal and hepatic side-effects are mandatory.

**15.6.3 Observation**

Observation until the development of clinically evident metastatic disease might represent a viable option for patients with a Gleason score ≤ 7, PSA recurrence longer than 2 years after surgery and a PSADT longer than 10 months. In these patients, the median actuarial time for the development of metastasis will be 8 years and median time from metastasis to death will be another 5 years.

**15.6.4 MANAGEMENT OF PSA RELAPSE AFTER RADICAL PROSTATECTOMY**

- Local recurrences are best treated by salvage radiation therapy with 64-66 Gy at a PSA serum level ≤ 1.5 ng/mL (grade B recommendation)
- Expectant management is an option for patients with presumed local recurrence unfit for, or unwilling to undergo, radiation therapy (grade B recommendation)
- PSA recurrence indicative of systemic relapse is best treated by early ADT resulting in decreased frequency of clinical metastases (grade B recommendation)
- LHRH analogues/orchiectomy or bicalutamide at 150 mg/day can both be used when there is indication for hormonal therapy (grade A recommendation)

**15.7 Management of PSA failures after radiation therapy**

In a recent review of the data of the Cancer of the Prostate Strategic Urologic Research Endeavor (CaPSURE) comprising 2,336 patients with CaP, Grossfeld et al. [64] demonstrated that 92% of patients initially irradiated received ADT for secondary treatment of PSA progression. In the absence of salvage procedures, the mean time interval from biochemical to clinical progression is approximately 3 years. Therapeutic options in these patients are ADT or local procedures, such as salvage radical prostatectomy, cryotherapy and interstitial radiation therapy (65-70). Salvage radical retropubic prostatectomy, however, has not gained widespread acceptance due to its associated morbidity, namely incontinence, local recurrences and rectal injuries. However, in well-selected patients, salvage radical retropubic prostatectomy might result in long-term disease-free survival. One has to consider that most series reporting on salvage radical prostatectomy included patients who were treated in the pre-PSA era without modern radiotherapeutic techniques and local recurrences were usually detected at a late stage. Therefore, complications associated with the procedure were quite high with up to 65% of the patients suffering from treatment-related morbidities. Up to 60% of the patients planned for salvage radical prostatectomy had to undergo anterior or total exenteration for locally extensive disease associated with a high rate of local recurrences and a mean time to progression of only 1.3 years (45-48).

Recent reports analyzing patients being operated upon in the last decade have described far more optimistic outcomes after salvage radical prostatectomy. In the series of Gheiler et al. (70), 40 patients with a mean PSA of 14 ng/mL underwent salvage radical prostatectomy. When stratified by PSA less than or greater than...
10 ng/mL, the 3-year disease-specific survival was 68% and 26%, respectively. In the series reported by Garzotto and Wajsman (69), 24 patients underwent radical cystoprostatectomy or radical prostatectomy with neoadjuvant ADT. Neoadjuvant ADT was associated with a lower rate of positive surgical margins (21%) as compared to patients in whom androgen deprivation failed and who exhibited a positive surgical margin rate of 80%. The authors demonstrated that the disease-specific survival strongly correlated with the surgical margin status. At a mean follow-up of 5 years the disease-specific survival rate was 95% and 44% for those with negative and positive surgical margins, respectively. Vaidya and Soloway (71) demonstrated a low complications rate, good postoperative continence and only one biochemical recurrence 36 months after salvage radical prostatectomy. Similar data have been achieved by Stephenson et al. (72), who reported on 100 consecutive patients undergoing radical salvage prostatectomy associated with a very low rate of perioperative complications. The 5-year progression-free rates have improved and the results are similar to those of standard radical prostatectomy in cases of similar pathological stages. The 10-year cancer specific and overall survival rates are in the range of 70% to 75% and 60% to 66% in contemporary series. In most contemporary series, organ-confined disease, negative surgical margins and the absence of seminal vesicle and/or lymph node metastases are favourable prognosticators associated with a better disease-free survival of approximately 70-80% as compared to 40-60% in patients with locally advanced CaP.

In general, salvage radical retropubic prostatectomy should only be considered in patients with a low comorbidity, a life expectancy of at least 10 years, an organ-confined CaP ≤ T2, Gleason grade < 7, and presurgical PSA < 10 ng/mL. In all other patients, accurate presurgical staging is not easily defined after radiation therapy, increasing the risk not only for the necessity of anterior and total extirpation procedures but also for associated complications and decreased long-term disease-specific survival.

15.7.1 Salvage cryosurgical ablation of the prostate (CSAP) for radiation failures
Salvage cryosurgery has been proposed as an alternative to salvage prostatectomy with the potential advantage of less morbidity and equal efficacy. There are only very few studies available and the results are not very promising. Pfisters et al. (73) report on 150 patients having undergone CSAP for PSA recurrences following radiotherapy (n = 110) or other extensive pre-treatment (n = 40). After a mean follow-up of 13.5 months, 58% of patients exhibited biochemical failure and only 31% demonstrated undetectable PSA serum levels. The complications associated with salvage CSAP were significant and occurred in basically all patients, with the main complications being urinary incontinence (73%), obstructive symptoms (67%), impotence (72%) and severe perineal pain (8%). After a 1-year follow-up, incontinence resolved in the majority of patients with a persistent significant incontinence in 22% of the patients (53%). According to a recent study by Cespedes et al. (74), the risk for urinary incontinence and impotence at least 12 months following CSAP are as high as 28% and 90%, respectively. In addition, 8% to 40% of the patients complained about persistent rectal pain and an additional 4% of men have undergone surgical procedures for the management of treatment-associated complications.

15.7.2 Salvage brachytherapy for radiation failures
The experience with salvage brachytherapy for radiation failures is very limited and there is only one study that includes a representative number of patients and a mean follow-up of 64 months (75-78).

Grado et al. (77) treated 49 patients with transperineal TRUS-guided brachytherapy and reported 3- and 5-year disease-free survival rates of 48% and 43%, respectively. Beyer et al. (78) reported a 5-year biochemical freedom from relapse in 34% to 53% of patients, with local cancer control achieved in 98% of patients. However, the complication rate was quite severe with 27% of the patients becoming incontinent, 14% having the need for a palliative TURP due to acute urinary retention and another 4% and 2% suffering from rectal ulcers and permanent colostomy.

15.7.3 Observation
Patients with signs of local recurrence only (low-risk patients with late recurrence and a slow PSA rise) and who are not opting for second-line curative options are best managed by observation alone. A retrospective cohort analysis of hormonal therapy versus watchful waiting in 248 men with PSA failure after radiotherapy showed no advantage for hormonal therapy in the subgroup of men with a PSADT of ≥12 months after radiotherapy. The 5-year metastasis-free survival rate was 88% with hormonal therapy vs 92% with watchful waiting (p=0.74) (79).

15.7.4 MANAGEMENT OF PSA RELAPSE AFTER RADIATION THERAPY
- Local recurrences may be treated by salvage radical prostatectomy in carefully selected patients (grade C recommendation)
- CSAP and interstitial brachytherapy are alternative experimental procedures in patients not suitable for surgery (grade C recommendation)
- ADT is an option in patients with patients with presumed systemic relapse (grade B recommendation).
### GUIDELINES FOR SECOND-LINE THERAPY AFTER CURATIVE TREATMENT

1. **Presumed local failure after radical prostatectomy**
   - Patients with presumed local failure only may be candidates for salvage radiotherapy. This should be given with at least 64 Gy and preferably before PSA has risen above 1.5 ng/mL. Other patients are best offered a period of watchful waiting (active monitoring) with possible hormonal therapy later on (grade B recommendation).

2. **Presumed local failure after radiotherapy**
   - Selected patients may be candidates for salvage radical prostatectomy although patients should be informed concerning the comparatively high risk of complications. Other patients are best offered a period of watchful waiting (active monitoring) with possible hormonal therapy later on (grade C recommendation).

3. **Presumed distant +/- local failure**
   - There is some evidence that early hormonal therapy may be of benefit in delaying progression and possibly achieve a survival benefit in comparison with delayed therapy. The results are not without controversy. Local therapy is not recommended except for palliative reasons (grade B recommendation).

### REFERENCES


   11590813

   12756084

   7538500


   15491116

   14767288


90 UPDATE MARCH 2005
16 HORMONE-REFRACTORY PROSTATE CANCER (HRPC)

16.1 Background
Cancer of the prostate is a heterogeneous disease and our understanding of the mechanism of androgen independence remains incomplete (1-5). Androgen ablation provides a selective advantage to androgen-independent cells that grow and eventually comprise the majority of the tumour. An alteration in normal androgen signalling probably has a central role in the pathogenesis of androgen-independent CaP. Androgen independence may be mediated through mutations of the androgen receptor gene that alter expression of the androgen receptor or its sensitivity to androgens (3-5). The fact that androgen receptor mutations are found in only a subpopulation of cells in the tumour suggests that these changes alone are unlikely to account fully for the entire spectrum of the androgen-independent state.

Many studies have focused on the deregulation of apoptosis in the development of androgen-independent disease. High levels of bcl-2 expression are seen with greater frequency as CaP progress, and a mechanism whereby bcl-2 induces its antiapoptotic effect may be the regulation of microtubule integrity (7-9). The fact that the most active chemotherapeutics in hormone-refractory prostate cancer (HRPC) work by inhibiting microtubule formation suggests that these findings may be clinically relevant. The tumour suppressor gene p53 is more frequently mutated in androgen-independent CaP. Over-expression of bcl-2 and p53 in prostatectomy specimens have been shown to predict an aggressive clinical course (10-12).

Peptide growth factors may have an important role in the progression of CaP. Epidermal growth factor is a potent mitogen of prostate stromal and epithelial cells. It is produced in high levels locally and acts as a paracrine stimulator. In androgen-independent tumours, autocrine stimulation may become more important, which, with epidermal growth factor, could allow unregulated growth (13).

16.2 Definition of HRPC
Hormone-refractory prostate cancer is a very heterogeneous disease including a variety of different patient cohorts with significant different median survival times (Table 13). Many different terms have been used to describe cancers that relapse after initial hormonal ablation therapy, including HRPC, androgen-independent cancers and hormone-independent cancers (1).

Table 13: Estimated natural mean survival of patients with HRPC presenting with different clinical scenarios.

<table>
<thead>
<tr>
<th>Patient characteristics</th>
<th>Estimated mean survival</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asymptomatic PSA †</td>
<td></td>
</tr>
<tr>
<td>• No metastases</td>
<td>18 – 20 months</td>
</tr>
<tr>
<td>• Minimal metastases</td>
<td>14 months</td>
</tr>
<tr>
<td>• Extensive metastases</td>
<td>9 – 12 months</td>
</tr>
<tr>
<td>Symptomatic PSA †</td>
<td></td>
</tr>
<tr>
<td>• Minimal metastases</td>
<td>9 months</td>
</tr>
<tr>
<td>• Extensive metastases</td>
<td>6 – 8 months</td>
</tr>
</tbody>
</table>
The precise definition of recurrent or relapsed CaP remains controversial. Recently, the various groups have published practical recommendations that should be adhered to when defining HRPC (14-16). Androgen-independent, but hormone-sensitive CaP, has to be differentiated from true HRPC from the outset. Whereas the first group still responds to secondary hormonal manipulations, such as antiandrogen withdrawal, oestrogens and corticosteroids, the latter is resistant to all hormonal measures.

**Definition of HRPC**

1. Serum castration levels of testosterone.
2. Three consecutive rises of PSA 2 weeks apart resulting in two 50% increases over the nadir.
3. Antiandrogen withdrawal for at least 4 weeks*.
4. PSA progression despite secondary hormonal manipulations*.
5. Progression of osseous or soft tissue lesions (3).

* Either antiandrogen withdrawal or one secondary hormonal manipulation should have been done in order to fulfil the criteria for HRPC.

**16.3 Assessing outcome of treatment in androgen-independent CaP**

In general, therapeutic outcome should be assessed according to the guidelines for the evaluation of response to treatment in solid tumours recently published by the RECIST group (Response Evaluation Criteria In Solid Tumours) (19). However, 80-90% of patients do not have bidimensionally measurable disease. Patients who have cancers with primarily soft tissue disease frequently have a different prognosis to those who have only osseous metastases. Osteoblastic bone metastases remain difficult to quantify accurately. There remains no general agreement regarding the methodology of measuring response (20-25). Determination of the cause of death in CaP patients is often unreliable, suggesting that overall, rather than disease-specific, survival rate may be a more valid endpoint (24).

Many contemporary studies use PSA as a marker of response, although there is no general consensus on what the magnitude and duration of decline in PSA level should be. The greatest use of PSA in this context is as a rapid screening tool to test new agents for activity. However, conflicting evidence is emerging regarding the role of PSA as a marker for response, and wide fluctuations have been seen in PSA values, indicating a transient effect of drugs on PSA production. Therefore, knowledge of the effects of a drug on PSA expression is the key to interpreting PSA response data, which must be viewed together with other clinical data (25-32). Despite these considerations, it has been reproducibly shown that ≥ 50% PSA decline in pre-treatment PSA following therapy is associated with a significant survival advantage (26,33). Kelly et al. (26) reported a statistically significant survival advantage in 110 patients if they had ≥ 50% PSA decline as opposed to those who did not (8.6 months vs > 25 months, respectively). Likewise, Smith et al. (33) demonstrated an increase in survival if a PSA decline ≥ 50% was maintained for at least 8 weeks, resulting in a mean survival time of 91 weeks as compared to only 38 weeks in those without this decrease. Molecular markers are just beginning to be evaluated. In a promising study, positive findings using reverse transcriptase-polymerase chain reaction (RT-PCR) were correlated with poor survival (34); however, these data have to be corroborated in other trials before recommendations can be made with regard to their clinical use.

In patients with symptomatic osseous lesions, pain reduction or complete pain relief might also be used as parameters to assess palliative therapeutic response (35).

Growing numbers of investigators advocate subjective endpoints. Since a significant survival benefit from chemotherapy in HRPC has not just been demonstrated in a subset of patients, the success of treatment may rely on redefining the goals of therapy (15,24,36). Currently, investigators should rely on clearly defined endpoints in trials that are sufficiently powered to answer the question posed, report each response parameter individually (rather than as a complete or partial response), use PSA response only in conjunction with other clinical parameters of response and consider QoL endpoints in symptomatic patients.

**16.4 RECOMMENDATIONS FOR ASSESSING THERAPEUTIC RESPONSE**

- PSA decline ≥ 50% maintained for 8 weeks is associated with a significantly better outcome as compared to a PSA decline < 50% (level of evidence:1a)
- In non-osseous metastases from HRPC, assessment should adhere to the RECIST criteria (level of evidence:1b)
- In patients with advanced symptomatic metastatic HRPC, therapeutic response can be assessed best by improvement of symptoms (level of evidence: 1b).

**16.5 Androgen deprivation in androgen-independent CaP**

Androgen-independent CaP implies that disease progression occurs despite castration. Therefore, castration levels of testosterone must first be documented. A serum testosterone level < 20 to 50 ng/mL should be documented at
16.6 Secondary hormonal therapy
For the patient with progressive disease after androgen deprivation, multiple therapeutic options are available and include antiandrogen withdrawal, addition of antiandrogens, oestrogenic compounds, adrenolytic agents and novel approaches (41). The therapeutic algorithm given in Figure 1 summarizes the various treatment modalities and the responses to be expected.

Figure 1: Flowsheet of the potential therapeutic options after PSA progression following initial hormonal therapy.

<table>
<thead>
<tr>
<th>PSA ↓ &gt;50%</th>
<th>Mean Duration of Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>100%</td>
<td></td>
</tr>
<tr>
<td>LHRH-analogues</td>
<td>Metastatic Prostate Cancer</td>
</tr>
<tr>
<td>subcapsular orchiectomy</td>
<td>36 months</td>
</tr>
<tr>
<td>CAB or anti-androgen monotherapy</td>
<td></td>
</tr>
<tr>
<td>60-80%</td>
<td></td>
</tr>
<tr>
<td>Addition of Anti-Androgens</td>
<td>4-6 months</td>
</tr>
<tr>
<td>Addition of Anti-Androgens or LHRH-analogues</td>
<td></td>
</tr>
<tr>
<td>25-40%</td>
<td></td>
</tr>
<tr>
<td>Substitution of flutamide with high-dose bicalutamide</td>
<td>4-6 months</td>
</tr>
<tr>
<td>30-40%</td>
<td></td>
</tr>
<tr>
<td>Anti-Androgen Withdrawal</td>
<td>5-6 months</td>
</tr>
<tr>
<td>40-60%</td>
<td></td>
</tr>
<tr>
<td>Secondary Hormonal Manipulation such as adrenal testosterone inhibitors, low-dose DES, steroids</td>
<td>4-8 months</td>
</tr>
<tr>
<td>50-70%</td>
<td></td>
</tr>
<tr>
<td>Non-hormonal Therapy such as chemotherapy</td>
<td>10-12 months</td>
</tr>
</tbody>
</table>

16.7 Antiandrogen withdrawal syndrome
In 1993, Kelly and Scher (42) reported clinical and PSA responses in men who discontinued flutamide therapy upon development of progressive disease. The antiandrogen withdrawal syndrome was a critical discovery in terms of understanding the biology of androgen independence, interpreting clinical trials and treating patients (42-46). Approximately one-third of patients will respond to antiandrogen withdrawal as indicated by a ≥ 50% PSA decrease with a median duration of response of approximately 4 months (Table 14). Antiandrogen withdrawal responses have also been reported after treatment with bicalutamide and megestrol acetate (48-53). The availability and more favourable toxicity profile of secondary hormonal therapies allow the clinician to consider these drugs for the growing category of asymptomatic patients for whom chemotherapy is difficult to justify, but who, due to increasing serum PSA level, want treatment outside clinical trials. However, observation remains a viable choice for asymptomatic patients.
Table 14: Frequency and duration of PSA response following antiandrogen withdrawal.

<table>
<thead>
<tr>
<th>Antiandrogen</th>
<th>N</th>
<th>&gt; 50% PSA</th>
<th>Duration (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flutamide</td>
<td>57</td>
<td>16 (28%)</td>
<td>4.0</td>
</tr>
<tr>
<td>Flutamide</td>
<td>82</td>
<td>12 (15%)</td>
<td>3.5</td>
</tr>
<tr>
<td>Flutamide</td>
<td>39</td>
<td>11 (28%)</td>
<td>3.7</td>
</tr>
<tr>
<td>Flutamide</td>
<td>21</td>
<td>7 (33%)</td>
<td>3.7</td>
</tr>
<tr>
<td>Bicalutamide</td>
<td>17</td>
<td>5 (29%)</td>
<td>5.0</td>
</tr>
</tbody>
</table>

16.8 Treatment alternatives after initial hormonal therapy

Except in patients with non-castration testosterone levels, it remains difficult to predict which subset of individuals is most likely to respond to secondary hormonal strategies (46). Bicalutamide is a non-steroidal antiandrogen that demonstrates a dose response, which means that 200 mg of bicalutamide normalizes PSA more effectively than 50 mg of bicalutamide in patients with androgen-dependent CaP (54). The benefits of adding an antiandrogen, such as bicalutamide or flutamide, to gonadal suppression at the time of PSA failure seems to result in PSA declines in a minority of patients only (55-57).

Approximately 10% of circulating androgen in humans is secreted by the adrenal glands. In androgen-independent states, some tumour cells must retain sensitivity to androgens, as a further decrease in circulating androgen levels by bilateral adrenalectomy or drugs that inhibit adrenal steroidogenesis can induce a clinical response. Aminoglutethimide, ketoconazole and corticosteroid act primarily via this mechanism (58-62) resulting in a PSA response in about 25% of patients treated lasting for about 4 months. The simultaneous addition of ketoconazole to antiandrogen withdrawal, however, results in a significantly increased PSA response (32% vs 11%) and a longer time to PSA progression (8.6 vs 5.9 months) compared to antiandrogen withdrawal alone (62), as has been documented in a recent, prospective, randomized phase III trial including 260 patients with androgen-independent CaP.

CaP normally expresses oestrogen receptors, which are upregulated after androgen ablation in animal models. In-vitro oestrogens can activate mutant androgen receptors that have been isolated in androgen-independent CaP. Anti-oestrogens in CaP have been reported to have measurable response rates of only 0–10% (63). Alternatively, high-dose oestrogens have been reported to have salvage objective responses. The mechanism for the effect has been postulated to be due to the mitotic arrest of direct cytotoxic effects on the cells, perhaps through an apoptotic mechanism (64,65). Recently, DES has been evaluated in two studies (66,67) including 21 and 32 patients. A positive PSA response was achieved in 43% and 80%, respectively; the estimated survival at 2 years was 63%. However, even at low doses, 31% of the patients developed deep venous thrombosis and 7% experienced myocardial infarction. In another prospective randomized phase II trial, the clinical efficacy of the herbal supplement PC-SPES and DES were tested in a cohort of 90 patients with PSA progression following initial androgen deprivation (68). A PSA decline ≥ 50% with was noted in 40% with PC-SPES and in 24% with DES. Median time to progression was 5.5 months with PC-SPES and 2.9 months with DES; the differences were statistically not significant.

16.9 Non-hormonal therapy (cytotoxic agents)

Based on prospective randomized clinical phase III trials, several proven chemotherapeutic options are available for the management of HRPC with metastatic disease (Table 15). In two recent phase III trials, a significant improvement in median survival of approximately 2 months could be demonstrated for docetaxel-based chemotherapy as compared to a combination of mitoxantrone and prednisone (69,70). In the TAX 327 study (70), 1,006 patients with metastatic HRPC were randomly assigned to mitoxantrone at 12 mg/m² every 3 weeks, docetaxel at 75 mg/m² every 3 weeks, or docetaxel at 30 mg/m² weekly for 5 of every 6 weeks. The median survival was 16.5 months in the mitoxantrone group and 18.9 months (p < 0.001) and 17.4 months in the docetaxel 75 mg/m² every 3 weeks and docetaxel 30 mg/m² for 5 of every 6 weeks, respectively. A ≥ 50% PSA decline was achieved in 45% and 48% of men in the docetaxel-treated groups compared to 32% in the mitoxantrone group (p < 0.001). Significant pain reduction was achieved in 22% of the patients in the mitoxantrone group compared to 35% (p=0.01) and 31% (p=0.08) in the docetaxel-treated groups. Adverse events were similar among the different treatment groups. However, QoL was significantly improved in both docetaxel-treated groups.

In the SWOG 99-16 trial (69), 674 patients with metastatic HRPC were randomly assigned to receive mitoxantrone at 12 mg/m² every 3 weeks or docetaxel and estramustine at 60 mg/m² every 3 weeks. In an intention to treat analysis, the median survival was 17.5 months and 15.6 months (p=0.02) in the docetaxel and the mitoxantrone groups, respectively. Also, the median time to progression was significantly longer in the docetaxel group with 6.3 months compared with 3.2 months in the mitoxantrone group (p < 0.001). A PSA decline of ≥ 50% was achieved in 50% and 27% patients of the docetaxel and the mitoxantrone group, respectively. Pain relief was similar among both groups, though side-effects occurred significantly more often in the docetaxel group.
Table 15: PSA response rates, mean survival and time to progression, and pain reduction in the large prospective randomized phase III trials documenting clinical efficacy of chemotherapy in patients with HRPC.

<table>
<thead>
<tr>
<th>Study</th>
<th>n</th>
<th>PSA ≥ 50%</th>
<th>pain</th>
<th>survival</th>
<th>TTP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tax 327</td>
<td></td>
<td>32%</td>
<td>22%</td>
<td>16.5 months</td>
<td>--</td>
</tr>
<tr>
<td>Docetaxel, 75mg/m²</td>
<td>45%</td>
<td>35%</td>
<td>18.9 months</td>
<td>--</td>
<td></td>
</tr>
<tr>
<td>Docetaxel, 30mg/m²</td>
<td>48%</td>
<td>31%</td>
<td>17.4 months</td>
<td>--</td>
<td></td>
</tr>
<tr>
<td>SWOG 99-16</td>
<td></td>
<td>50%</td>
<td>17.5 months</td>
<td>6.3 months</td>
<td></td>
</tr>
<tr>
<td>Docetaxel/EMP</td>
<td>338</td>
<td>27%</td>
<td>15.6 months</td>
<td>3.2 months</td>
<td></td>
</tr>
<tr>
<td>CALGB 9182</td>
<td></td>
<td>38%</td>
<td>--</td>
<td>12.3 months</td>
<td>2.3 months</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>123</td>
<td>22%</td>
<td>--</td>
<td>12.6 months</td>
<td>3.7 months</td>
</tr>
<tr>
<td>Mitoxantrone/HC</td>
<td>119</td>
<td>22%</td>
<td>--</td>
<td>12.6 months</td>
<td>3.7 months</td>
</tr>
<tr>
<td>Tannock et al.</td>
<td></td>
<td>33%</td>
<td>29%</td>
<td>43 weeks</td>
<td>18 weeks</td>
</tr>
</tbody>
</table>

TTP = median time to progression; EMP = estramustine; HC = hydrocortisone; Pred = prednisone.

Despite these encouraging results, the time point to initiate cytotoxic regime in patients with HRPC remains controversial. Although, it appears evident that chemotherapy should be started in patients with metastatic HRPC, there are no data available with regard to the therapeutic efficacy of early chemotherapy in patients with PSA rise only. There, at least, exists the recommendation that two consecutive increases in PSA over a previous reference value should exist and that the PSA level should exceed 5 ng/mL. Therefore, the indication for the initiation of chemotherapeutic regimes has to made on an individual basis.

In order to further improve treatment results, some experimental phase I and phase II trials combining taxanes with anti-bcl-2, calcitriol, exisulid, and thalidomide are underway resulting in a PSA response of about 60% (71-74). In a randomized phase II trial of docetaxel plus thalidomide (71), 75 men with chemotherapy-naïve HRPC were randomized to receive either docetaxel at 30 mg/m² for 5 of every 6 weeks or docetaxel at the same dose and schedule plus thalidomide at 200 mg orally each day. A PSA decline of ≥ 50% was higher in the combination-treated group (53%) compared to the docetaxel-alone treated group (37%) without reaching statistical significance. Median progression-free survival and overall survival were 5.9 months and 68% at 18 months as compared to 3.7 months and 43% in the docetaxel-alone group without reaching statistical significance. However, side-effects were significant, with thromboembolic events occurring in 28% of the combination arm as compared to no such events in the docetaxel arm.

Mitoxantrone with corticosteroids (30,75) has been extensively studied primarily in patients with symptomatic osseous lesions due to HRPC. In the CALGB 9182 study (75), 244 patients with symptomatic metastatic HRPC were randomized to either receive mitoxantrone plus hydrocortisone at 12 mg/m² every 3 weeks or to hydrocortisone alone. Although no differences were observed with regard to survival, PSA response, and median time to progression, QoL was significantly improved in the combination arm. In the other trial (30), 161 men with painful osseous metastases due to HRPC were randomized to either receive mitoxantrone plus hydrocortisone at 12 mg/m² every 3 weeks or to hydrocortisone alone. Although no differences were observed in terms of pain reduction as compared to prednisone alone (12%, p=0.01); furthermore, duration of palliation was longer in patients who received mitoxantrone (43 vs 18 weeks, p < 0.0001). There were no significant differences with regard to PSA response and median survival time. Although none of the studies demonstrated any survival benefit for the patients, QoL was improved significantly due to pain reduction.

Alternative treatments evaluated in prospective clinical phase II trials, including pegylated doxorubicin, a combination of paclitaxel, carboplatin and estramustine, a combination of vinblastine, doxorubicin and radonuclides and a combination of docetaxel and mitoxantrone, have demonstrated encouraging results (76-81). The lack of representative randomized phase III trials and still unknown efficacy in terms of long-term outcome are the major problems associated with all of these studies.

The synergy observed for estramustine in combination with other drugs that target microtubule action has generated promising results in prospective clinical trials (82). Estramustine plus vinblastine has been the most studied estramustine combination; although different doses of estramustine and vinblastine have been used in prospective randomized trials, significant PSA and measurable responses have been reported in three separate studies. Although time to progression and frequency of ≥ 50% PSA decrease was significantly higher in the combination arm, median survival did not differ significantly between the estramustine and the estramustine plus vinblastine arms.

UPDATE MARCH 2005
Intravenous cyclophosphamide has been tested in multiple trials. Current interest has focused on oral cyclophosphamide, which appears to be less toxic than when given intravenously and may have greater activity (83,84). A study of the combination of oral cyclophosphamide and oral etoposide in 20 patients was similarly encouraging (83). Cisplatin and carboplatin have activity against CaP as single agents, but their synergy with etoposide or paclitaxel in vitro and in the treatment of other diseases, such as lung and ovarian cancer, is well documented. As estramustine is also synergistic with these drugs, combinations of three agents are now being tested. A combination of estramustine, etoposide and cisplatin (or carboplatin) has significant activity against poorly differentiated HRPC. A combination of estramustine, etoposide and paclitaxel has also recently been reported to produce high response rates (77).

Suramin activity against HRPC is likely to be mediated through the inhibition of binding of growth factors, such as transforming growth factor b, to their receptors. Although the ultimate role of suramin in the treatment of HRPC is still undetermined, recent results renew some of this agent’s initial promise (78-80).

16.10 Palliative therapeutic options

The majority of patients with HRPC have painful bone metastases. The two beta-emitting radioisotopes, strontium-89 and samarium-153, can partially or completely decrease bone pain in up to 70% of patients. Early use can make subsequent administration of chemotherapy more difficult because of myelosuppression (81,84). Critical issues of palliation must be addressed while considering additional systemic treatment, including management of pain, constipation, anorexia, nausea, fatigue and depression, which frequently occur (i.e. palliative external beam radiation, cortisone, analgesics and antiemetics).

Common complications due to skeletal metastases include bone pain, vertebral collapse or deformity, pathologic fractures and spinal cord compression. Recently, the use of bisphosphonates to inhibit osteoclast-mediated bone resorption and activity of osteoclast precursors has demonstrated a clinically significant effect in terms of prevention of skeletal complications and reduction of pain, or even total pain relief, in patients with HRPC. In the largest single phase III trial (88), 643 men with HRPC metastatic to the bone were randomized to receive zoleodronic acid at 8mg or 4mg every 3 weeks for 15 consecutive months or placebo. At 15 months and at 24 months of follow-up, there was a significant reduction in skeletal-related events in the zoleodronic acid treated group as compared to the placebo group (44% vs 33%, p=0.021). The frequency of pathological fractures was significantly lower in the zoleodronic acid group compared with the placebo group (13.1% vs 22.1%, p=0.015). Furthermore, the time to first skeletal-related event was significantly prolonged in the zoleodronate group thereby significantly improving QoL. Currently, bisphosphonates could be proposed to patients with HRPC bone metastases in order to prevent skeletal complications.

Pain due to osseous metastases is one of the most debilitating complications of HRPC. Bisphosphonates have been proven to be highly effective with a response rate of 70-80%, which, associated with a low frequency of side-effects, makes bisphosphonates to be an ideal medication for palliative therapy of advanced HRPC (35,89). Bisphosphonates should be considered early in the management of symptomatic HRPC.

Hormone refractory CaP is usually a debilitating disease, often affecting the elderly male. A multidisciplinary approach is required with input from medical oncologists, radiation oncologists, urologists, nurses and social workers (90).

16.11 SUMMARY ON TREATMENT AFTER HORMONAL THERAPY

- It is recommended to cease antiandrogen therapy once PSA progression is documented (grade B recommendation)
- Four to six weeks after discontinuation of flutamide or bicalutamide, an eventual anti-androgen withdrawal (AAW) effect will become apparent (grade B recommendation)
- No clear cut recommendation can be made regarding the most effective drug for secondary hormonal manipulations since data from randomized trials are scarce (grade C recommendation).
16.12 GUIDELINES AND RECOMMENDATIONS FOR CYTOTOXIC THERAPY IN HRPC
1. In patients with a PSA rise only, 2 consecutive increases of PSA serum levels above a previous reference level should be documented (grade B recommendation)
2. Prior to treatment, PSA serum levels should be > 5ng/mL to assure correct interpretation of therapeutic efficacy (grade B recommendation)
3. Potential benefits of cytotoxic therapy and expected side effects should be discussed with each individual patient (grade C recommendation)
4. In patients with metastatic HRPCA, and who are candidates for cytotoxic therapy, docetaxel at 75 mg/m² every 3 weeks has shown a significant survival benefit (grade A recommendation)
5. In patients with symptomatic osseous metastases due to HRPCA either docetaxel or mitoxantrone with prednisone or hydrocortisone are viable therapeutic options (grade A recommendation).

16.13 GUIDELINES FOR PALLIATIVE MANAGEMENT OF HRPC
1. Patients with symptomatic and extensive osseous metastases cannot benefit from medical treatment with regard to prolongation of life
2. Management of these patients has to be directed at improvement of QoL and mainly pain reduction
3. Effective medical management with the highest efficacy and a low frequency of side-effects represents the major goal.

16.14 RECOMMENDATIONS FOR PALLIATIVE MANAGEMENT OF HRPC
1. Bisphosphonates may be offered to patients with skeletal metastases (mainly zoledronic acid has been studied) to prevent osseous complications (grade A recommendation)
2. Palliative treatments such as radionuclids, external beam radiotherapy, adequate use of analgesics should be considered early on in the management of painful osseous metastases (grade B recommendation).

16.15 REFERENCES


ABBREVIATIONS USED IN THE TEXT

AAW  antiandrogen withdrawal effect
ADT  androgen deprivation therapy
ASAP  atypical small acinar proliferation
ASTRO American Society of Therapeutic Radiology and Oncology
bNED  actuarial biochemical freedom of disease/biochemical non-evidence of disease
CAB  complete androgen blockade
CaP  cancer of the prostate
CaPSURE  Cancer of the Prostate Strategic Urologic Research Endeavor
CPA  cyproterone acetate
3D-CRT  three-dimensional conformal radiation therapy
CSAP  cryosurgical ablation of the prostate
CT  computed tomography
DES  diethylstilboestrol
DHT  dihydrotestosterone
digital rectal examination
EBM  evidence-based medicine
ECOG  Eastern Cooperative Oncology Group
EORTC  European Organization for Research and Treatment of Cancer
ER-β  oestrogen receptor-beta
ERSPC  European Randomized Screening for Prostate Cancer
EU  European Union
FSH  follicle-stimulating hormone
FNCLCC  Fédération Nationale des Centres de Lutte Contre le Cancer
HIFU  high-intensity focused ultrasound
HRPC (HRPC)  hormone-refractory prostate cancer
HRQoL  health-related quality of life
HT  hormonal therapy
IAB  intermittent androgen blockade
IAD  intermittent androgen deprivation
ICI  intracavernosal injection
IMRT  intensity modulated radiotherapy
IPPS  International Prostatic Symptom Score
LH  luteinizing hormone
LHRH  luteinizing hormone-releasing hormone
LTAD  long-term androgen deprivation (ablation)
MRC  Medical Research Council
MRI  magnetic resonance imaging
NCI  National Cancer Institute
NHT  neoadjuvant hormonal therapy
NIH  National Institutes of Health
PAP  prostatic acid phosphatase
PET  positron emission tomography
PIN  prostatic intraepithelial neoplasia
PLCO  Prostate, Lung, Colorectal, Ovary trial
PSA  prostate-specific antigen
PSADT  prostate-specific antigen doubling time
PSMA mRNA  prostate specific membrane antigen for messenger RNA
QoL  quality of life
QUALYs  quality of life adjusted gain in life years
RECIST group  Response Evaluation Criteria In Solid Tumours group
RITA  radiofrequency interstitial tumour ablation
RTOG  Radiation Therapy Oncology Group
RT-PCR  reverse transcriptase-polymerase chain reaction
SEER (database)  Surveillance, Epidemiology, and End Results database of the National Cancer Institute (USA)
STAD  short-term androgen deprivation (ablation)
TNM  Tumour Node Metastasis (classification)
TRUS  transrectal ultrasonography
TURP  transurethral resection of the prostate

UPDATE MARCH 2005
Levels of evidence

<table>
<thead>
<tr>
<th>Level</th>
<th>Type of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>Evidence obtained from meta-analysis of randomized trials</td>
</tr>
<tr>
<td>1b</td>
<td>Evidence obtained from at least one randomized trial</td>
</tr>
<tr>
<td>2a</td>
<td>Evidence obtained from one well-designed controlled study without randomization</td>
</tr>
<tr>
<td>2b</td>
<td>Evidence obtained from at least one other type of well-designed quasi-experimental study</td>
</tr>
<tr>
<td>3</td>
<td>Evidence obtained from well-designed non-experimental studies, such as comparative studies, correlation studies and case reports</td>
</tr>
<tr>
<td>4</td>
<td>Evidence obtained from expert committee reports or opinions or clinical experience of respected authorities</td>
</tr>
</tbody>
</table>


Grades of guideline recommendations

<table>
<thead>
<tr>
<th>Grade</th>
<th>Nature of recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Based on clinical studies of good quality and consistency addressing the specific recommendations and including at least one randomized trial</td>
</tr>
<tr>
<td>B</td>
<td>Based on well-conducted clinical studies, but without randomized clinical trials</td>
</tr>
<tr>
<td>C</td>
<td>Made despite the absence of directly applicable clinical studies of good quality</td>
</tr>
</tbody>
</table>