Antiandrogen therapy immediately after surgery for prostate cancer improved survival


**Question**
In men with node-positive prostate cancer, is survival prolonged by starting antiandrogen therapy immediately after surgery rather than delaying it until the appearance of disease progression?

**Design**
Randomized [allocation concealed*†, unblinded,* controlled trial with median 7.1-year follow-up.

**Setting**
6 U.S. university centers.

**Patients**
98 men (median age 66 y) who had clinically localized prostate cancer (≤ stage T2), had had radical prostatectomy and bilateral pelvic lymphadenectomy within the past 12 weeks, and had histologically confirmed nodal metastases.

**Intervention**
Patients were allocated to antiandrogen therapy with either goserelin, 3.6 mg subcutaneously every 28 days, or bilateral orchiectomy (n = 47) or to be followed until the appearance of disease progression other than newly detectable or rising levels of prostate-specific antigen (PSA) (n = 51).

**Main Outcome Measures**
Death, death from prostate cancer, and recurrence.

**Main Results**
Analysis was by intention to treat. At a median follow-up of 7.1 years, more men in the observation group had died from any cause (hazard ratio [HR] 3.0, 95% CI 1.2 to 7.3; P = 0.02) or from prostate cancer (HR 6.2, CI 1.8 to 21.5; P < 0.01) than had men who received immediate antiandrogen therapy (Table). Disease recurrence was greater in the observation group than in the immediate antiandrogen therapy group (HR 12.2, CI 5.1 to 29.1; P < 0.001) (Table). Baseline characteristics or type of antiandrogen therapy did not affect survival.

**Conclusion**
In men with node-positive prostate cancer, starting antiandrogen therapy immediately after surgery rather than delaying it until the appearance of disease progression improved survival.

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*See Glossary.
†Information provided by author.

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Immediate therapy</th>
<th>Observation</th>
<th>RRR (95% CI)</th>
<th>NNT (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death</td>
<td>15%</td>
<td>35%</td>
<td>58% (12 to 81)</td>
<td>5 (3 to 34)</td>
</tr>
<tr>
<td>Death from prostate cancer</td>
<td>6.4%</td>
<td>31%</td>
<td>80% (40 to 93)</td>
<td>4 (3 to 10)</td>
</tr>
<tr>
<td>Any recurrence</td>
<td>15%</td>
<td>82%</td>
<td>82% (66 to 91)</td>
<td>2 (2 to 2)</td>
</tr>
</tbody>
</table>

*Abbreviations defined in Glossary; RRR, NNT, and CI calculated from data in article.

**Commentary**
The study by Messing and colleagues convincingly shows better outcomes for men who, after radical prostatectomy and pelvic lymphadenectomy, are found to have lymph node metastases and are immediately treated with androgen deprivation rather than waiting for disease progression to appear. Such men are at high risk for disease progression, even those with low PSA levels after surgery alone (only 16% of the control group in this study showed no evidence of disease progression over 7 years). These results are consistent with a trial done in the United Kingdom that also showed improved cancer-specific survival for men with locally advanced or asymptomatic metastatic disease who were treated with immediate androgen deprivation (1).

One problem with long trials is that as new technologies appear, the trials’ results may be less helpful in practice than originally hoped. Currently, widespread PSA screening has resulted in a “stage shift,” with a small percentage of radical prostatectomies now associated with positive nodes.

Most men who are started on androgen deprivation show evidence of residual or recurrent cancer after surgery or radiation therapy solely on the basis of PSA levels. Those with recurrent disease have a better prognosis than node-positive patients, with a median time to a positive bone scan of 8 years without treatment (2). As a result, competing risks become more of an issue for patients with recurrent disease than for node-positive patients. Furthermore, the cumulative effects on health after years or decades of androgen deprivation are only now emerging. These consequences were reported only superficially in this study. Further trials comparing early with delayed androgen deprivation are needed before assuming that early androgen deprivation has a similar risk-benefit tradeoff in men who have either early evidence of disease recurrence or prognostic factors at the time of primary treatment that suggest a major risk for future recurrence.

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**References**