Identification of Intermediate Cell Types by Keratin Expression in the Developing Human Prostate

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BACKGROUND. The secretory acini of the adult human prostate contain basal, luminal, and intermediate types of exocrine cells. Intermediate cells are thought to play an important role in normal growth and neoplastic transformation. In this study we investigated whether this cell type is present in early stages of prostate development, using keratin antibodies specific for them.

METHODS. Autopic tissue from 11 prepubertal and 5 normal adult prostates was immuno-histochemically stained with four keratin antibodies capable of specifically detecting basal, luminal, or intermediate cell types.

RESULTS. Morphologically, in fetal prostate cells differentiation was often not evident. However, basally located cells usually displayed a basal-cell keratin-phenotype. Morphologically similar cells with more luminal localization expressed keratins typical of luminal cells, or of intermediate cells.

CONCLUSIONS. 1) In early stages of prostate development, cells with intermediate keratin-phenotype can be identified. 2) Their large numbers comply with a hierarchical pathway of cellular differentiation from basal to luminal cells. 3) The presence of intermediate cells at such an early fetal age may reflect their regulatory function in prostate development.


KEY WORDS: intermediate filament; immunohistochemistry; embryology; maturation; benign prostatic hyperplasia; etiology

INTRODUCTION

Although benign prostatic hyperplasia (BPH) is one of the most common conditions in elderly men [1], our knowledge concerning its etiology is fragmentary. The correlation between adult neoplastic growth and fetal prostate development was first hypothesized by McNeal [2]. He emphasized that BPH occurring later in life is the result of a “reawakening” of the embryonic capability in the adult, i.e., of the stromal inductive potential on epithelial growth [2]. Prostatic early development and morphogenesis are dependent on inductive signals originating in the mesenchymal compartment [3]. Furthermore, adult urogenital epithelia maintain the capability to respond to stromal mediators of growth and differentiation, and several peptide growth factors have been implicated in the development of BPH [4,5]. These studies on the embryology and development of the prostate provide important clues which may elucidate mechanisms of prostatic neoplasia.

In characterizing epithelial cell differentiation, antibodies against keratin (K) components of the cytoskeleton can be instrumental, as the 20 different members of this family are expressed in specific combinations in various types of epithelia. Moreover, their
expression is related to specific stages of differentiation (reviewed in Moll et al. [6], Ramaekers et al. [7], and Nagle [8]). Previous keratin expression studies in the fetal prostate showed that of 30–36-week of gestation, only basal-cell keratins are expressed. In the 7-month neonate and 1-year-old infant, distinct changes in keratin expression occur, and a population of cells expressing basal-type keratins is found along with a population expressing luminal-type keratins [9]. However, data concerning the exact keratin-phenotype of these cells and possible transitions in keratin expression during fetal development are not available. At present, the keratin-phenotype for prostate exocrine epithelial cells comprises, among others, keratins 5 and 14, also called complex keratins found in basal cells, while luminal cells contain, among others, keratins 8 and 18, called simple keratins [10,11]. It is thought that during differentiation, basal cells acquire simple keratins (K8 + K18) and lose their basal-cell keratins (K5 + K14).

Based on light and electron microscopic studies in humans and dogs, a third cell type has been identified, having characteristics of both basal and luminal cells [12,13]. In the postnatal rat prostate these cells contain both basal-cell keratins 5 and 15 and luminal-cell keratins 8 and 18 [14]. Bonkhoff et al. [15] observed a subset of basal cells which coexpress basal-cell keratins and PSA (prostatic-specific antigen is usually only expressed in luminal cells). This is considered evidence for the presence of a group of cells with intermediate differentiation [15]. Our own studies confirm the presence of an intermediate cell type in the secretory acini of the normal adult human prostate, in BPH, in prostate carcinoma (PCa), and in involution-regeneration processes in rat prostate [16–18]. We demonstrated that these cells contain keratin 18 and express the basal-keratin antibody RCK103. Based on these observations we proposed that this cell type could be the same as the amplifying cells described in the stem-cell model [19].

Because knowledge concerning intermediate cells in the developing human prostate is incomplete, we investigated the keratin-phenotype of epithelial cells in the prostate during various stages of human development to see whether cells with an intermediate cell keratin-phenotype could be identified. This may provide more information regarding the similarities between the process of prostate maturation and development of BPH.

### MATERIALS AND METHODS

#### Tissues

Tissue specimens used in this study comprised 11 prepuberal prostates taken during autopsy in 5 fetuses (17, 19, 27, 32, and 38 weeks of gestation), 3 infants (2, 3.5, and 7 months old), and 3 prepubertal males (1 year and 7 months, 1 year and 7 months, and 11 years old). For comparative purposes, prostate tissue from 5 adults (27–49 years old) was also used. Tissue blocks were fixed in 10% phosphate-buffered formalin and processed through paraffin. All patients died of non-prostatically related diseases, and none of them had been hormonally treated. Postmortem delay did not exceed 10 hr (5.6 ± 2.4 hr). From each tissue block at least five consecutive sections were cut. The tissue block in the fetal prostate was a transversal section through the entire prostate. In the adults, a representative section was taken without identifying the anatomic location.

#### Antibodies

Four monoclonal antibodies were used in this study and are summarized in Table I.

Briefly, RCK102 directed against K5 and K8 is a broadly crossreacting keratin antibody. RCK102 stains most epithelial tissues, while nonepithelial tissues do not react with this antibody. We used RCK102 as a general epithelial marker [20].

Monoclonal antibody 34BE12 recognizes K1, K5, K10, and K14. As K1 and K10 are characteristic of

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Specificity</th>
<th>Localization in prostatic tissue</th>
<th>Dilution</th>
<th>Source</th>
<th>Reference</th>
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</thead>
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<tr>
<td>RCK102</td>
<td>K5 and K8</td>
<td>All epithelial cells</td>
<td>1:10</td>
<td>Dr. G. van Muijen, Nijmegen, Netherlands</td>
<td>20</td>
</tr>
<tr>
<td>34BE12</td>
<td>K1, K5, K10, and K14</td>
<td>Basal cells</td>
<td>1:50</td>
<td>DAKO, Carpinteria, CA</td>
<td>21</td>
</tr>
<tr>
<td>RCK103</td>
<td>K5, etc.</td>
<td>Basal and some luminal cells</td>
<td>1:5</td>
<td>Dr. G. van Muijen, Nijmegen, Netherlands</td>
<td>17,18</td>
</tr>
<tr>
<td>CK18 (clone DC-10)</td>
<td>K18</td>
<td>Luminal cells</td>
<td>1:40</td>
<td>DAKO, Glostrup, Denmark</td>
<td>16,22</td>
</tr>
</tbody>
</table>
stratified keratinizing squamous epithelium, which is not found in the prostate, this antibody recognizes K5 and K14 in prostatic tissue [21].

RCK103 recognizes K5 and a number of other keratins that are not yet characterized. In the prostate it reacts with basal cells and intermediate cells, and to identify these last cells they must coexpress K18 [16,17].

CK18 (clone DC-10) recognizes K18. In the prostate this keratin is found in luminal cells, for which it is considered a marker [22].

### Immunohistochemistry

Four-micrometer tissue sections were cut from paraffin blocks, mounted on coated slides, and dried overnight. After deparaffination, antigen retrieval was performed: for CK18 (DC-10), microwave heating was performed according to the manufacturer’s instructions. For the antibodies RCK102 and RCK103, antigen retrieval was achieved according to a modified protocol [23] by submerging the slides in a 0.5% periodic acid solution and microwave heating for two cycles of 5 min each (180 W). For 34\beta E12, slides were incubated with 0.1% Pronase XIV solution (P-5147, Sigma Chemical Co.) dissolved in 10 mM Tris, 1 mM EDTA, pH 7.5, at 37°C for 10 min. Sections were incubated with the primary antibodies at 4°C overnight. Biotinylated secondary antibody and AB complex were used (Elite Vector ABC kit, Amersham, Burlingame, CA). Peroxidase activity was visualized with diaminobenzidine (DAB) as a chromogen. Sections were briefly counterstained in hematoxylin and coverslipped.

### Evaluation of Keratin Immunostaining Results

The number of positively staining cells in the entire section was estimated for the four keratin-antibodies semiquantitatively and independently by two of us (Y.X., F.S.). Any discrepancies were discussed and consensus was reached in all cases. The number of positive staining cells was estimated and recorded as follows: −, no cells stained; +/−, incidental positive cells, <1%; +, between 1–25% of cells; ++, between 26–50% of cells; ++++, between 51–75% of cells; and +++++, between 76–100% of cells. Staining intensity
was not separately graded, as it was often heterogeneous within a specimen and between specimens. The pattern of staining, however, was constant. Staining was evaluated separately for the three cell types in the budding tips, also referred to as the distal compartment (see Results), and for the more proximal tubules referred to as the ductal compartments which are directly adjacent to the urethral orifice and which show immature glandular lumens. Positive controls consisted of prostate tissues from surgery specimens, which are known to react strongly with the antisera used [17]. Bovine serum albumin was used instead of the primary antibodies for negative controls.

RESULTS

Light Microscopy

Topographically, prostate epithelial cells could be subdivided into three types, and are illustrated schematically in Table II. First, basally located cells were directly adjacent to the basal membrane, which separated them from the stromal compartment. These cells had no direct contact with glandular lumen. Second, intermediately located cells were found above the basal cells and underneath the luminally located cells which lined the prostate acinus. In early development, many layers of these suprabasal cells were seen in solid cords in the peripheral zone of prenatal and infantile prostate. Third, luminally located cells lined the acini. Under them, basally located cells and sometimes intermediately located cells were found. In early development, it was not easy to identify these cells separately.

The epithelial compartment of the 17- and 19-week fetuses consisted of budding solid cords of cells in the peripheral parts of the gland, surrounded by highly cellular stroma (Fig. 1a). The basement membrane was inconspicuous. Cells in the solid cords had relatively large ovoid nuclei and scant cytoplasm. Columnar cells were not recognizable. Luminal differentiation was found in the proximal parts of the tubules. The luminal portion of these ducts was lined by nonciliated low columnar or cuboid cells with underlying multilayered basaloid cells. In the 27-week fetus and in the older fetuses, the branching process started to appear in the peripheral parts of the prostate. It was characterized by increased numbers of solid buds and tubules lined by epithelium, similar to those found in the younger fetus. The ducts in the vicinity of the urethra were sometimes lined by squamous metaplastic epithelium (Fig. 2c).

The prostates from the 4 infants showed a histologic picture different from that of the fetuses. The number of developing solid cords had increased and also the number of ducts. The stroma was a little less cellular. The basally located cells were the same as in the fetus, but more cords had lumina lined by columnar cells.

In the two prostates from children, 16 and 19 months old, solid cords of epithelial cells were mostly replaced by arborizing tubules lined by one or more
Fig. 2. Immunoperoxidase staining pattern of developing (a,c,e,g) and adult (b,d,f,h) prostate after staining with RCK102 (a,b), 34βE12 (c,d), RCK103 (e,f), and CK18 (g,h). Tissue sections were from prostates of (a) 19-week fetus, (c) 32-week fetus, (e) 11-year-old male, and (g) 32-week fetus. Arrow indicates squamous metaplastic epithelium (c). Original magnification: objective ×40.
<table>
<thead>
<tr>
<th>Age</th>
<th>Site</th>
<th>RCK102</th>
<th>34βE12</th>
<th>RCK103</th>
<th>CK18</th>
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<tbody>
<tr>
<td>17 weeks</td>
<td>B</td>
<td>NA</td>
<td></td>
<td>NA</td>
<td></td>
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<tr>
<td>19 weeks</td>
<td>B</td>
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<tr>
<td>27 weeks</td>
<td>B</td>
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<tr>
<td>32 weeks</td>
<td>B</td>
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<tr>
<td>38 weeks</td>
<td>B</td>
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<tr>
<td>2 months</td>
<td>B</td>
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<td>3.5 months</td>
<td>B</td>
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<tr>
<td>7 months</td>
<td>B</td>
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<tr>
<td>1 year, 4 months</td>
<td>B</td>
<td></td>
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<tr>
<td>1 year, 7 months</td>
<td>B</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>11 years</td>
<td>B</td>
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*TABLE III. Cellular Keratin Phenotype on Budding Tips of Developing Acini and Mature Glands in the Prostate*
layers of small cells with relatively large nuclei and scant cytoplasm. Luminal low columnar or cuboidal cells, usually with large central nuclei, were also seen. Polarity and secretory vacuoles in the supranuclear portion were hardly discernible, and often it was difficult to distinguish these cells from adjacent cells with more basaloid features. The prostate of the prepubertal 11-year-old male basically showed the same picture as in the infants, especially in the distal segment of the ductal acini. This was characterized by a more elaborate formation of arborizing irregularly-shaped glands consisting of a single layer of small spindle-shaped basal cells underlying a single layer of tall columnar luminal cells with periapical vacuolation of their cytoplasm. Stroma was still predominant but less cellular in comparison to younger prostates (Fig. 1b). For comparison, the histology of adult prostate acini is shown in Figure 1c.

**Keratin Immunophenotyping**

Keratin expression was separately evaluated in the distal segment of branching glandular structures or budding tips, and in the proximal ductal compartment associated with the urethra, for the basally-located, intermediately-located, and luminally-located cells.

**Keratin Immunophenotyping in the Developing Prostate (Table III, Fig. 2)**

**Basally-located cells.** In 9 of 11 developing prostates, all basally-located cells were positive for RCK102 and 34βE12 (Fig. 2a,c). In 2 cases about half of the basally-located cells stained for 34βE12. Monoclonal antibody RCK103 was intensely immunoreactive with basally located cells in both budding tips and ducts of the developing prostate (Fig. 2e). However, there was some variability in the number of positively staining cells, and in 3 cases approximately 50% of cells stained, and one case was negative. CK18 was not expressed in the basal-cell compartment of any of the cases (Fig. 2g).

**Intermediately-located cells.** In all but 2 cases, all intermediately located cells stained with the RCK102 antibody (Fig. 2a). 34βE12, a marker for basal cells, stained 50–75% of cells in the intermediate compartment of 10 cases (Fig. 2c), while in one case only sporadic cells stained. RCK103 stained the cells in this compartment in most cases. However, the percentage of cells staining was considerably lower than that found in the basal compartment (Fig. 2e). In one case, only sporadic cells stained. CK18 was detected in 10 cases in the intermediate compartment. One case was negative. Immunoreactivity was variable, with 4 cases showing positivity in 50% of cells, 2 cases showing positivity in 25% of cells, and 4 cases showing only sporadic staining (Fig. 2g).

**Luminally-located cells.** RCK102 was positive in all cases in most cells (Fig. 2a). In 9 cases, between 25–50% of luminal cells stained with 34βE12 (Fig. 2c). In

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**TABLE III. Continued**

<table>
<thead>
<tr>
<th>Age</th>
<th>Site</th>
<th>RCK102</th>
<th>34βE12</th>
<th>RCK103</th>
<th>CK18</th>
</tr>
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<tbody>
<tr>
<td>Mature gland</td>
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<td></td>
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<tr>
<td>27 years</td>
<td>B</td>
<td></td>
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<tr>
<td>39 years</td>
<td>B</td>
<td></td>
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<tr>
<td>42 years</td>
<td>B</td>
<td></td>
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<tr>
<td>48 years</td>
<td>B</td>
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<tr>
<td>49 years</td>
<td>B</td>
<td></td>
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</table>

*A completely solid bar indicates that between 75–100% of cells stained. A ¼ solid bar indicates that between 50–75% of cells stained. A ½ solid bar indicates that between 25–50% of cells stained. A ¾ solid bar indicates that between 1–25% of cells stained. An open bar indicates that no cells stained. Horizontal lines mean incidental positive in <1% of cells. B, basally located cells; I, intermediately located cells; L, luminally located cells; NA, not available; F, fetus. For details of scoring, see Materials and Methods.*
one case, only sporadic cells were immunoreactive, and in one case the luminal compartment was negative. The cases with the least reactivity were the infant and the prepubertal prostate. In most cases, RCK103 showed the same or slightly less immunoreactivity than with intermediately-located cells (Fig. 2e). In one case, only sporadic cells stained, and one other case was completely negative. Keratin 18 was detectable in the luminal-located cells of all cases. However, the percentage of positive cells varied. In 6 cases, virtually all cells in the luminal compartment stained (Fig. 2g), in 3 cases, 75%, in one case, 50%, and in one other case, only 25%. For K18, there seemed to be a trend toward lower levels of immunoreactivity in luminal cells with lower fetal age.

**Keratin Immunoreactivity in Ductal Compartments**

In general, ductal compartments showed immunoreactivity that paralleled that of the budding tips. Only salient features of keratin phenotyping are described. Differential expression of the various keratin antibodies between basal and luminal cells was often more obvious. Most striking was the focal expression of RCK103 in the luminal cell compartment, with 34βE12 showing positivity in 2 cases. In the fetuses of 17 and 27 weeks of gestation, ductal structures were not observed in the slides.

**Keratin Immunophenotyping in Adult Prostate**

Keratin expression in the adult prostate was the same as previously described in frozen sections [17]. Basal cells intensely expressed 34βE12 and RCK103, while CK18 was not expressed (Fig. 2d,f,h). Luminal cells only expressed CK18. However, 3 cases showed sporadic RCK103 positivity, and one case showed scanty expression of 34βE12. RCK102 was positive in almost all epithelial cells in the adult prostate (Fig. 2b).

**DISCUSSION**

Luminal and basal cell types encountered in the adult prostatic epithelium are two distinctly different cellular populations with regard to structure and function [24,25]. Early reports suggested that they originate independently [6,26,27]. More recent studies, however, provide evidence for the existence of a pluripotent progenitor cell type for luminal cells [28]. Ultrastructural studies on human, rat, and dog prostatic tissue suggest that basal cells are actively engaged in growth. They are not highly specialized in a structural sense but resemble undifferentiated secretory epithelial cells [12,13,24]. In the developing prostate, however, the typical morphologic basal-cell and luminal-cell features could not be readily identified.

Early studies on prostate development were based mainly on light and electron microscopic results. They showed that the prostatic buds appear at about week 12 of amenorrhea. The buds are formed from the mucosa of the urogenital sinus, which protrudes through the basement membrane to colonize the surrounding mesenchyme. Initially they are solid budding tips composed of monotonous cells with basaloid appearance. Ultrastructural studies show that at 13 weeks of gestation, luminal cells with dense secretion granules and nuclear polarization may be identified in the acinar lumen. The other cells present in these acini are less well-differentiated and have been hypothesized to represent the equivalents of basal cells in the adult prostate [29-31].

Antibodies to cytoskeletal proteins of the keratin subclasses have been used as a relatively new tool to study embryonic development, particularly with regard to differentiation [7,26, and references therein]. These studies show that changes in levels of differentiation are accompanied by distinct transitions in expression of individual keratins. In the present report we investigated whether application of a small panel of keratin antibodies could give a more profound insight into prostatic differentiation. And as such, this is an extensive report describing keratin expression in the fetal prostate. Due to the absence of keratin antibodies specific for intermediate cells, they are identified by their coexpression of antibodies RCK103 and CK18, and actually this MAb RCK103 is mainly found in the basal cell compartment. In a previous study we found that epithelial cells in the adult prostate could be characterized into basal cells (K14+/K5+/K18−), luminal cells (K14−/K5−/K18+), and intermediate cells (K14+ /K5−/K18+). We have suggested that these cells could well be the missing link between basal and luminal cell types during differentiation, and that they could play a role in neoplastic transformation in the adult prostate [17]. An interesting question is whether this cell type is present in the developing prostate, and if so when may it first be identified?

At 17 weeks of gestation, we observed variable differentiation-specific keratins in the immature prostate. Basally-located cells were positive for both 34βE12 and RCK103, but negative for CK18. Therefore, their keratin phenotype was K14+/K5+/K18−. This pattern is the same as that of spindle-shaped basal cells in adult prostates [17,18]. Most probably these basally-located cells are the same as adult prostate basal cells, which have proliferative potential [12,32].

The luminal-located cells were partially positive for 34βE12 and RCK103, and practically all contained K18. Therefore, their keratin phenotype is
K14+/K5+/K18+. These cells have a keratin phenotype different from that of their adult counterparts (K14+/K5+/K18+). This could be explained by contending that even though they are lumina1y located they are still not fully differentiated luminal cells, and that they still have basal-cell and intermediate-cell characteristics in terms of their keratin phenotype despite their luminal location, meaning that the morphologic transition of these cells, from basaloid to luminal, is preceded by a transition in keratin-phenotype. In some adult prostatic tissues we also noticed that sporadic cells in the luminal compartment had the basal-cell keratin-phenotype. This is consistent with previous reports and is probably associated with focal basal-cell hyperplasia [18,33]. Furthermore, comparing the intermediate cell type between the human adult prostate and the developing prostate, previous studies never found that intermediate cells express basal-cell keratin 34BE12 in the adult.

The intermediately-located cells displayed a heterogeneous keratin phenotype coexpressing both keratins found in basal and luminal cells (K14+/K5+/K18+). Furthermore, there are cells expressing basal keratins only and a number of cells expressing luminal-cell keratins only. It is possible that this intermingled pattern indicates the underlying gradual switch from basal cells with K14 and K5 phenotype to luminal cells with K18. This would strongly favor the role of these basal cells with K14 and K5 phenotype to luminal cells with K18. This would strongly favor the role of these intermediate cells that stemmed from basal cells and that will eventually differentiate into luminal cells. The simultaneous expression of both basal phenotypes in the fetal prostate reveals that the hierarchical expanding pathway from basal to luminal cells may be demonstrated in the most early stages of development. The fact that these cells are present in the fetus may indicate that they have a regulatory function in prostate development. Since the same intermediate cell type can be identified in the adult, this potential to transform probably remains throughout life. Pathologic stimulation, e.g., aging hormonal alteration, or disturbance in homeostasis (stromal-epithelial interactions), may reactivate this cell differentiation in terms of their keratin phenotype despite their luminal location, meaning that the morphologic transition of these cells, from basaloid to luminal, is preceded by a transition in keratin-phenotype. In some adult prostatic tissues we also noticed that sporadic cells in the luminal compartment had the basal-cell keratin-phenotype. This is consistent with previous reports and is probably associated with focal basal-cell hyperplasia [18,33]. Furthermore, comparing the intermediate cell type between the human adult prostate and the developing prostate, previous studies never found that intermediate cells express basal-cell keratin 34BE12 in the adult.

We conclude that in the fetal prostate, morphologic cell differentiation is preceded by differential keratin expression. In early stages of prostate development, cells with an intermediate keratin phenotype may be identified. The large numbers of these cells may reflect their position as a link between basal and luminal cells in a differentiation pathway.

ACKNOWLEDGMENTS

The authors thank Dr. E. Ruijter and Dr. E. Schaafsm for assistance in acquiring fetal prostate tissues, and Dr. E. Ruijter for critical comments in the preparation of this manuscript.

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