

# Expressed sequence tag profiling identifies developmental and anatomic partitioning of gene expression in the mouse prostate

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## Abstract

**Background:** The prostate gland is an organ with highly specialized functional attributes that serves to enhance the fertility of mammalian species. Much of the information pertaining to normal and pathological conditions affecting the prostate has been obtained through extensive developmental, biochemical and genetic analyses of rodent species. Although important insights can be obtained through detailed anatomical and histological assessments of mouse and rat models, further mechanistic explanations are greatly aided through studies of gene and protein expression.

**Results:** In this article we characterize the repertoire of genes expressed in the normal developing mouse prostate through the analysis of 50,562 expressed sequence tags derived from 14 mouse prostate cDNA libraries. Sequence assemblies and annotations identified 15,009 unique transcriptional units of which more than 600 represent high quality assemblies without corresponding annotations in public gene expression databases. Quantitative analyses demonstrate distinct anatomical and developmental partitioning of prostate gene expression. This finding may assist in the interpretation of comparative studies between human and mouse and guide the development of new transgenic murine disease models. The identification of several novel genes is reported, including a new member of the  $\beta$ -defensin gene family with prostate-restricted expression.

**Conclusions:** These findings suggest a potential role for the prostate as a defensive barrier for entry of pathogens into the genitourinary tract and, further, serve to emphasize the utility of the continued evaluation of transcriptomes from a diverse repertoire of tissues and cell types.

## Introduction

The normal function of the mammalian prostate gland is to enhance fertility by secreting buffers, proteins and protective agents that maintain sperm in a quiescent and intact state as they pass through the male and female reproductive tracts [1].

Much of the information pertaining to normal prostate physiology has been obtained through extensive developmental and biochemical analyses of the prostates of rodent species. Despite anatomical differences between rodents and humans, these studies have been instrumental for elucidating the

influence of androgens on prostate differentiation and growth, and for characterizing the protein and mineral constituents that comprise the unique prostate environment.

In humans, the prostate exhibits the distinctive attribute of sustained growth throughout life, a situation that contributes to both benign and malignant prostate pathologies [1]. Strikingly, the prevalence rates of benign prostate hypertrophy (BPH) and prostate carcinoma approach nearly 50% in American men by the age of 70 [2-4]. Despite extensive research efforts, the etiologies of these diseases remain poorly defined. In contrast to colon, skin and bladder epithelia, prostate epithelial cells are thought to be relatively better protected from environmental insults, and the cellular constituents exhibit low proliferation rates [5]. Androgenic hormones and hereditary factors influence both BPH and prostate carcinoma, but the specific mechanisms by which they alter cellular growth remain to be delineated.

As with other human health disorders, rodent models have been developed to aid in the scientific analysis of prostate diseases. Whilst early efforts focused extensively on the rat prostate [6,7] - in part due to the advantages of working with a relatively large gland - recent investigations have utilized the mouse increasingly, primarily as a result of the ease and power of manipulating the mouse genome [8]. The rodent prostate is comprised of four distinct lobes: ventral, anterior (also termed the coagulating gland), dorsal and lateral; the latter two are commonly grouped together and collectively referred to as the dorsolateral lobe [9]. These lobes are arranged circumferentially around the urethra and display characteristic patterns of ductal branching and secretory protein production. In contrast, the human prostate lacks a defined lobar architecture - it is organized in zones with distinct disease predispositions; carcinoma primarily develops in the peripheral zone and benign hypertrophy primarily occurs in the transition zone [10]. The anatomical and functional relationships between the rodent prostate lobes and the human prostate zones have not been definitively established, though it has been suggested that the human peripheral zone is most analogous to the rodent dorsolateral lobe based upon the observation that tumors induced in rodent prostates generally arise in these locations [7,11].

Although important insights pertaining to normal development and disease pathology can be obtained through detailed anatomical and histological assessments of mouse models, further mechanistic explanations are greatly aided through studies of gene and protein expression. In this context, the Cancer Genome Anatomy Project (CGAP) [12] and other large-scale sequencing efforts have sought to provide a comprehensive sequence and reagent set that encompasses genes expressed in diverse collections of human and mouse tissues [13]. However, a recent inventory of cDNA libraries and sequences archived in CGAP and the database of expressed sequence tags (ESTs) indicates that while 838 cDNA libraries

have been constructed from murine tissues and cell types, there is no mouse prostate representation (query 8.20.2003 in [12]). In this article we characterize the repertoire of genes expressed in the normal developing mouse prostate through the analysis of ESTs derived from mouse prostate cDNA libraries. The results of this analysis demonstrate distinct anatomical and developmental partitioning of gene expression, a finding that may assist in the interpretation of comparative studies between human and mouse, and further guide the evaluation and development of new transgenic murine disease models. The identification of several novel genes is reported, including a new member of the  $\beta$ -defensin gene family with prostate-restricted expression. This finding suggests a potential role for the prostate as a defensive barrier for entry of pathogens into the genitourinary tract.

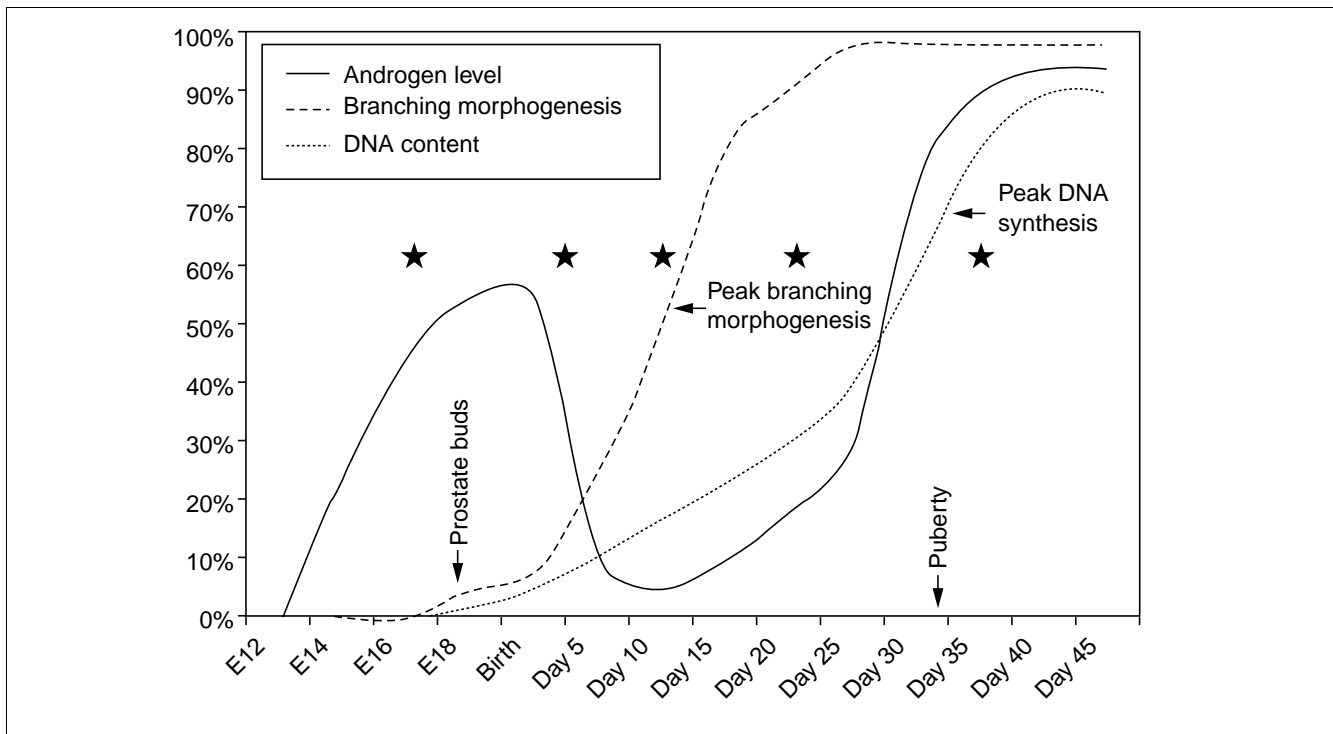
## Results

### Mouse prostate transcriptome

To assess the diversity of gene expression in the mouse prostate, we constructed and characterized cDNA libraries representing distinct stages of prostate development and maturation that began with the urogenital sinus (E16.5), and continued with prostates from two day old (neonatal), ten day old, 20 day old, 35 day old (puberty), three month old (normal adult), and 14 month old (aged adult) animals (Figure 1). A total of 62,046 ESTs were generated, of which 50,562 passed stringent quality assessments. Of these, 4,299 ESTs annotated to the mitochondrial genome. Assembly of the remaining prostate ESTs into distinct clusters or transcription units (TUs) was performed using the sequence assembly program Phrap [14] and each TU was assigned an annotation by BLAST comparisons against sequences in public databases. In accordance with definitions used by the FANTOM and RIKEN teams in assembling a mouse transcriptome using full-length cDNAs [15], we use the term TU to designate a cluster of transcripts or ESTs that contain a common core of sequence information transcribed from a segment of the genome. A TU may be defined by a single cDNA sequence or by multiple overlapping ESTs that share a common core sequence. Assembly and annotation of the 46,263 mouse prostate ESTs produced 15,009 unique TUs. Of these, 4,550 TUs are composed of more than one EST and 10,459 TUs are represented by a single EST. Of the 4,550 TUs that contain more than one EST, 3,567 are represented by sequences derived from more than one mouse prostate library. All sequences are archived for public accession in the mouse prostate expression database (mPEDB) [16,17].

### Gene expression alterations during mouse prostate development

To identify genes potentially involved with prostate development and maturation, we determined temporal expression changes by calculating transcript abundance levels in cDNA libraries constructed from different stages of prostate development. Pairwise comparisons of the datasets from each



**Figure 1**

Temporal events in mouse prostate development. Androgen levels (solid line) rise beginning at day E12, fall shortly after birth, and rise again at puberty [65]. Branching morphogenesis (dashed line) begins at approximately day E17 with prostate budding, peaks at approximately day 10 (after birth), and is essentially complete before puberty by day 35 [23]. Peak DNA synthesis (dotted line), representing the development of the prostate epithelial and stromal cell mass, occurs at approximately day 35 [66]. cDNA libraries were constructed from prostate tissues obtained at defined points of prostate development (stars).

timepoint were performed using Audic and Claverie's analysis method [18]. At a significance level of  $p = 0.001$ , 285 genes were differentially expressed between one or more time points (see Additional data file 1, Table 1). Applying the Bonferroni correction for multiple comparisons identified a cohort of 69 genes with highly significant differential expression between one or more developmental stages. Hierarchical clustering of these genes demonstrated distinct temporal partitioning of gene expression with notable cohorts increasing over time and others decreasing over time. In Figure 2, two clusters of genes are highlighted which show progressive increases or decreases in expression over time from urogenital sinus to adulthood. Several of these genes encode proteins involved in specialized prostate secretory activity such as spermine binding protein (*Sbp*) [19] and serine protease inhibitor Kazal type 3 (*Spink3*) [20]. Several are also known to be regulated by androgenic hormones, such as probasin (*Pbsn*), and increase in expression following the onset of puberty [21]. Other TUs with temporal changes in expression have no currently identified functional roles in prostate development.

### Gene expression compartmentalization between mouse prostate lobes

The rodent prostate gland is comprised of four distinct anatomical lobes: dorsal, lateral, ventral and anterior, each with unique ductal branching patterns and different responses to androgenic hormones (Figure 3) [22,23]. Studies of secretory products from rodent prostates have identified specific expression patterns that can be assigned to individual lobes [24-26]. To further explore the functional differences between mouse prostate lobes, we compared the profiles of genes expressed in each lobe and identified 34 genes that exhibited statistically significant differential expression. This group was reduced to 17 genes after correcting for multiple comparisons (Table 2). The expression patterns of these genes clearly support conclusions of prior anatomical and biochemical studies demonstrating functional heterogeneity between lobes of the gland. Genes highly expressed in the anterior prostate relative to the other lobes included onzin (alias placenta specific 8; *Plac8*), pancreatic family type A ribonuclease 1 (*RNase1*), and experimental autoimmune prostatitis antigen (*Eap1*; a putative prostate transglutaminase), as well as several uncharacterized transcripts. Spermine binding protein (*Sbp*), and serine protease Kazal type 3 (*Spink3*) are relatively highly expressed in the ventral lobe, while transcripts with increased expression in the

**Table 1****Characterized genes with enhanced prostate expression**

Symbol	UniGene description	Tissues	UniGene ID
<i>Syp2</i>	Seminal vesicle protein 2	Genitourinary; aorta and vein	Mm.1286
<i>Msmb</i>	$\beta$ -microseminoprotein	Genitourinary	Mm.2540
<i>Galgt2</i>	UDP-N-acetyl-alpha-D-galactosamine	Colon; embryo	Mm.2807
<i>Nkx3-1</i>	NK-3 transcription factor locus 1 ( <i>Drosophila</i> )	Trophoblast stem cell	Mm.3520
<i>Svs6</i>	Seminal vesicle secretion 6	Genitourinary	Mm.3787
<i>Sva</i>	Seminal vesicle antigen	Genitourinary	Mm.4119
<i>Pbsn</i>	Probasin	Genitourinary; bone	Mm.8034
<i>Defb2</i>	Defensin $\beta$ 2	Head; epididymis	Mm.41981
<i>Sbp</i>	Spermine binding protein	Tongue; genitourinary; stomach	Mm.46428
<i>B3galt1</i>	B 1 3-galactosyltransferase polypeptide 1	Medulla oblongata; ganglion	Mm.57041
<i>ErbB3</i>	Erythroblastic leukemia viral oncogene homolog 3	Tumor; inner ear	Mm.57112
<i>Hoxd12</i>	Homeo box D12	Embryo; forelimb	Mm.57124
-	11 kDa secreted protein precursor	Whole skin	Mm.71887
<i>Mllt7</i>	Mixed lineage-leukemia translocation to 7 homolog	Tumor; genitourinary; spleen	Mm.88827
<i>Smok2</i>	Sperm motility kinase 2	Lung	Mm.88851
<i>Svs7</i>	Seminal vesicle protein secretion 7	Genitourinary	Mm.99349
<i>Fhl3</i>	Four and a half LIM domains 3	Placenta	Mm.100241
<i>Pappa</i>	Pregnancy-associated plasma protein A	Parthenogenote; tumor; genitourinary	Mm.103481
<i>Svs3</i>	Seminal vesicle secretion 3	Genitourinary; adipose	Mm.118769
-	EST similar to Acrosomal vesicle protein 1	Genitourinary; thymus; placenta	Mm.118804
<i>Svs2</i>	Seminal vesicle protein secretion 2	Genitourinary; aorta and vein	Mm.143501
<i>Amd2</i>	S-adenosylmethionine decarboxylase 2	Blastocyst	Mm.195848
<i>Acac</i>	Acetyl-coenzyme A carboxylase	Whole brain; branchial arches	Mm.196688
<i>Muc10</i>	Mucin 10 submandibular gland salivary mucin	Salivary gland	Mm.200411
<i>Lamc2</i>	Laminin gamma2 chain	Inner ear	Mm.213197
<i>Birc1f</i>	Baculoviral IAP repeat-containing 1f	Lymph node; colon	Mm.218759
<i>Tnep1</i>	Thrombospondin N-terminal domain	Whole brain	Mm.221237

dorsolateral prostate include receptor activity modifying protein 2 (*Ramp2*), a transcript with similarity to acrosomal vesicle protein-1, and several uncharacterized transcripts. We determined the lobe-specific expression patterns of several transcripts using quantitative PCR (QPCR), and found that the EST and QPCR methods produced qualitatively similar results (Figure 4).

Wubah *et al.* have reported the cloning and tissue distribution of a transcript termed ventral prostate predominant 1 (*Vpp1*) that was identified through mRNA differential display between different mouse prostate lobes [27]. QPCR and virtual expression analysis confirmed that the expression of *Vpp1* is primarily localized in the ventral prostate, with lower but detectable levels of expression in the dorsolateral and anterior prostate lobes (Figure 4).

**Figure 2** (see following page)

Differential gene expression during mouse prostate development. Statistical analysis of transcript abundance levels in stages of mouse prostate development identified 69 differentially expressed genes ( $p = 0.001$ ). Shown are cohorts with progressing or regressing levels over time. The intensity scale represents the fold-change in expression normalized relative to the lowest abundance measurement.

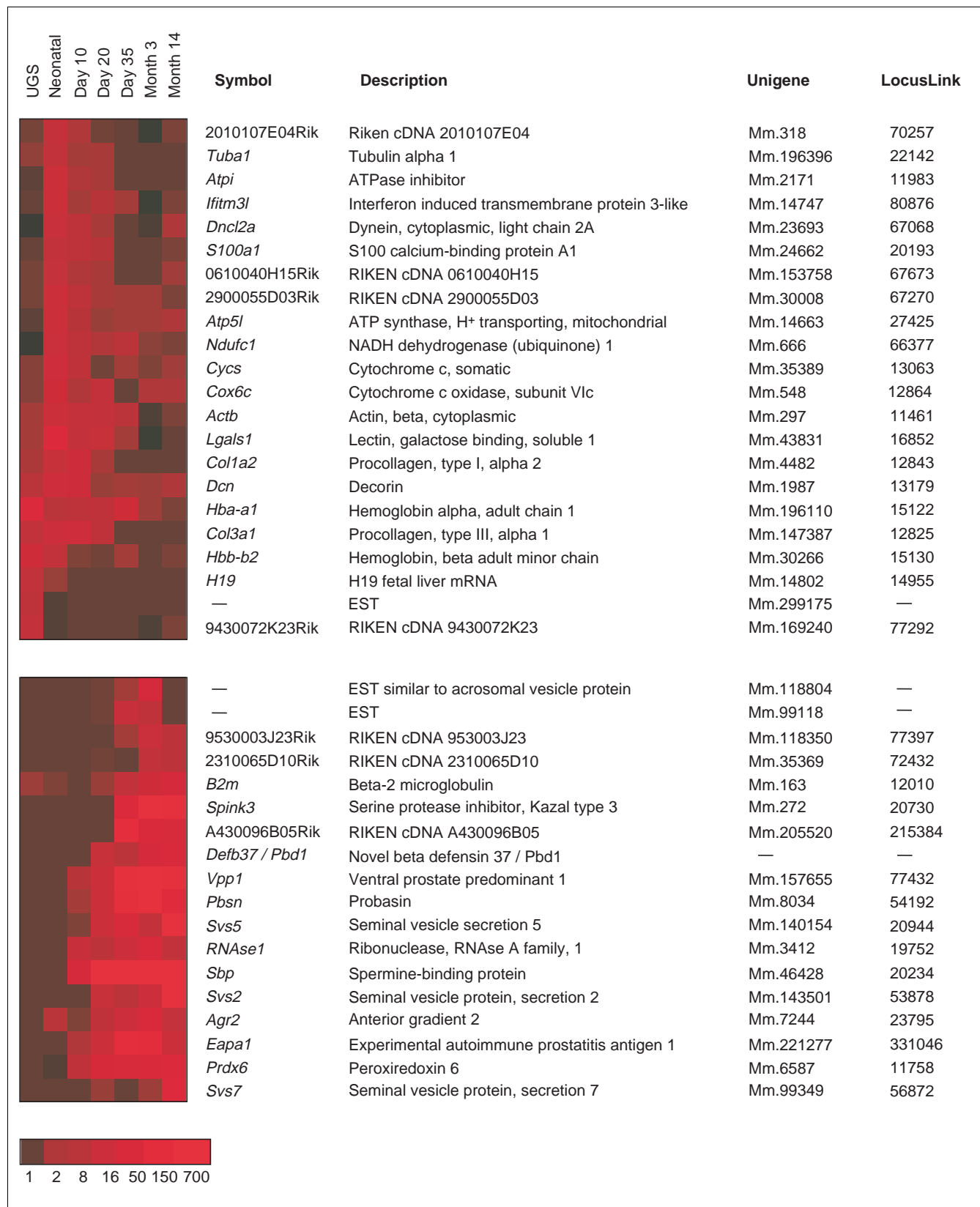
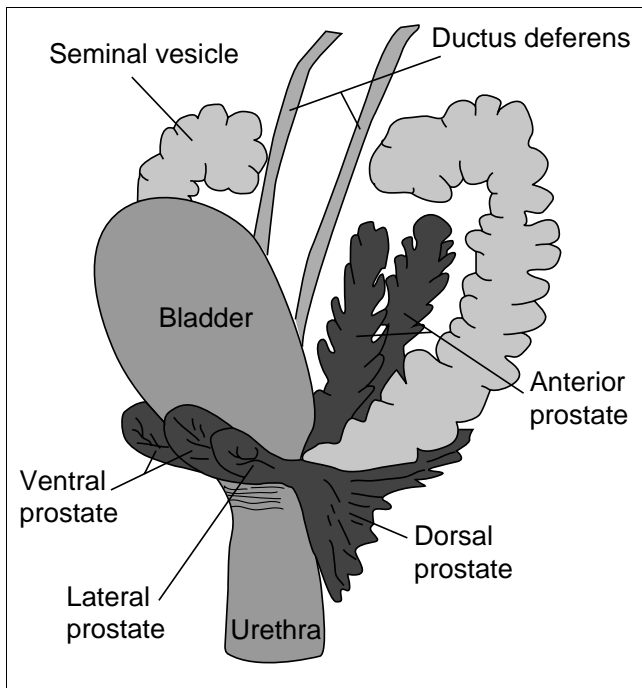


Figure 2 (see legend on previous page)



**Figure 3**  
A schematic diagram of the adult mouse genitourinary tract (lateral view), with the anatomically-distinct prostate lobes highlighted in gray. Reproduced with permission from [67,68].

Our EST results demonstrated that the probasin transcript was most highly expressed in the anterior prostate with lower - but still high - transcript abundance levels in the dorsolateral lobes, and minimal expression in the ventral lobe (Figure 4). QPCR quantitated the highest probasin expression in the dorsolateral lobes followed by the anterior prostate, with lowest levels in the ventral lobe. These results differ from studies of expression localization using the rat probasin promoter to drive transgene expression in mouse prostate epithelium where the highest levels were observed in the lateral and dorsal lobes, with lower expression in the ventral prostate and very low to absent expression in the anterior prostate [11,28]. Subsequent immunohistochemical and Western analyses with antibodies recognizing the mouse probasin protein demonstrated high levels of expression in the anterior as well as the dorsolateral mouse prostate lobes [29]. These results suggest that the rat probasin promoter may confer slightly different cellular specificity compared with the native mouse promoter, or that transgenic constructs alter the normal regional distribution of probasin expression.

#### Genes with expression enhanced or restricted to the mouse prostate

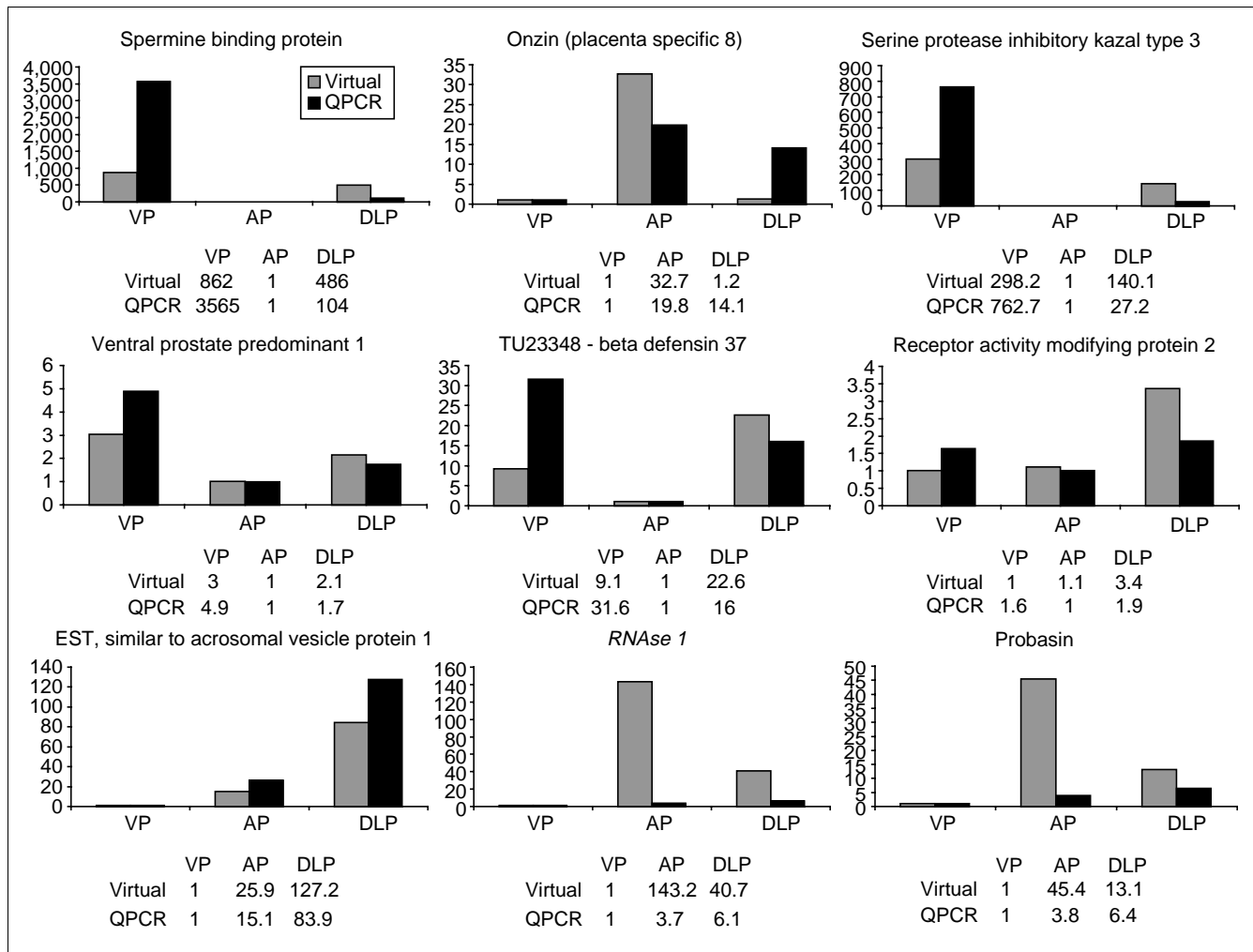
The comparative analysis of cDNA libraries representing the repertoire of genes expressed in the mouse prostate provided an opportunity to identify genes whose expression is enhanced or restricted to the mouse prostate relative to other normal tissues. The human prostate gland expresses several

transcripts and corresponding proteins in a highly tissue-restricted manner including prostate specific antigen (PSA) [30], human glandular kallikrein 2 (hK2) [31], prostase/ KLK4 [32] and prostate specific membrane antigen (PSMA) [33]. Studies of these proteins have provided insights into normal prostatic function, mechanisms of hormone-regulated gene expression and the development of prostate pathology. To identify transcripts preferentially expressed in the mouse prostate, we assigned each prostate TU to a UniGene cluster and then determined the tissue sources of all ESTs comprising the cluster. In total, 776 prostate TUs were mapped to UniGenes containing ESTs derived from three or fewer other tissues (see Additional data file 2). Only 28 of these TUs represent characterized genes (Table 1). Included among the 28 are NKX3.1 and probasin, both of which are known to have prostate-restricted patterns of expression [29,34]. Most of the prostate-enhanced TUs are represented by uncharacterized ESTs and full-length cDNAs sequenced by the RIKEN mouse transcript sequencing project [15]. Eighty TUs contained ESTs from at least two different prostate libraries. Of these, eight represent spliced gene products based on sequence comparisons with the mouse draft genome sequence. An additional 20 TUs had interesting features, such as high sequence conservation with human sequences, and may represent orthologous genes.

#### Identification of a gene encoding a putative prostate-specific $\beta$ -defensin

One TU, originally designated TU23348 after the original library clone identifier, was found to be expressed in multiple mouse prostate cDNAs libraries, lack representation in the UniGene or dbEST sequence databases, and exhibit sequence similarities to genes in the  $\beta$ -defensin family. We reasoned that since TUs specifically originating from the mouse prostate were not represented in the public sequence databases, TU23348 could represent a  $\beta$ -defensin with expression restricted to the prostate. To evaluate this possibility, we measured the relative abundance of TU23348 transcripts in multiple mouse tissues by QPCR. These results confirmed high levels of TU23348 expression in the prostate, with very low levels in testis and skeletal muscle, and no detectable transcripts in a wide range of other tissues (Figure 5a). A multiple tissue mRNA blot confirmed the RT-PCR findings (Figure 5b), and northern analysis identified a transcript size of approximately 0.4 kb. Northern analysis and QPCR also indicated that TU23348 expression is further compartmentalized within the mouse prostate having highest expression levels in the dorsolateral and ventral lobes, with very low to absent expression in the anterior prostate (Figure 5c).

A full-length TU23348 transcript was obtained through assemblies of mouse prostate cDNAs and by 5' and 3' rapid amplification of cDNA ends (RACE) reactions using mouse prostate cDNA as a template. The coding sequence comprises 219 nucleotides encoding a predicted polypeptide of 73 amino acids (Figure 6a). The nucleotide sequence comprises a 72 nt



**Figure 4**  
Comparative analysis of fold change of gene expression in adult prostate lobes using quantitative PCR (QPCR) (dark grey bars) and virtual EST measurements (light grey bars).

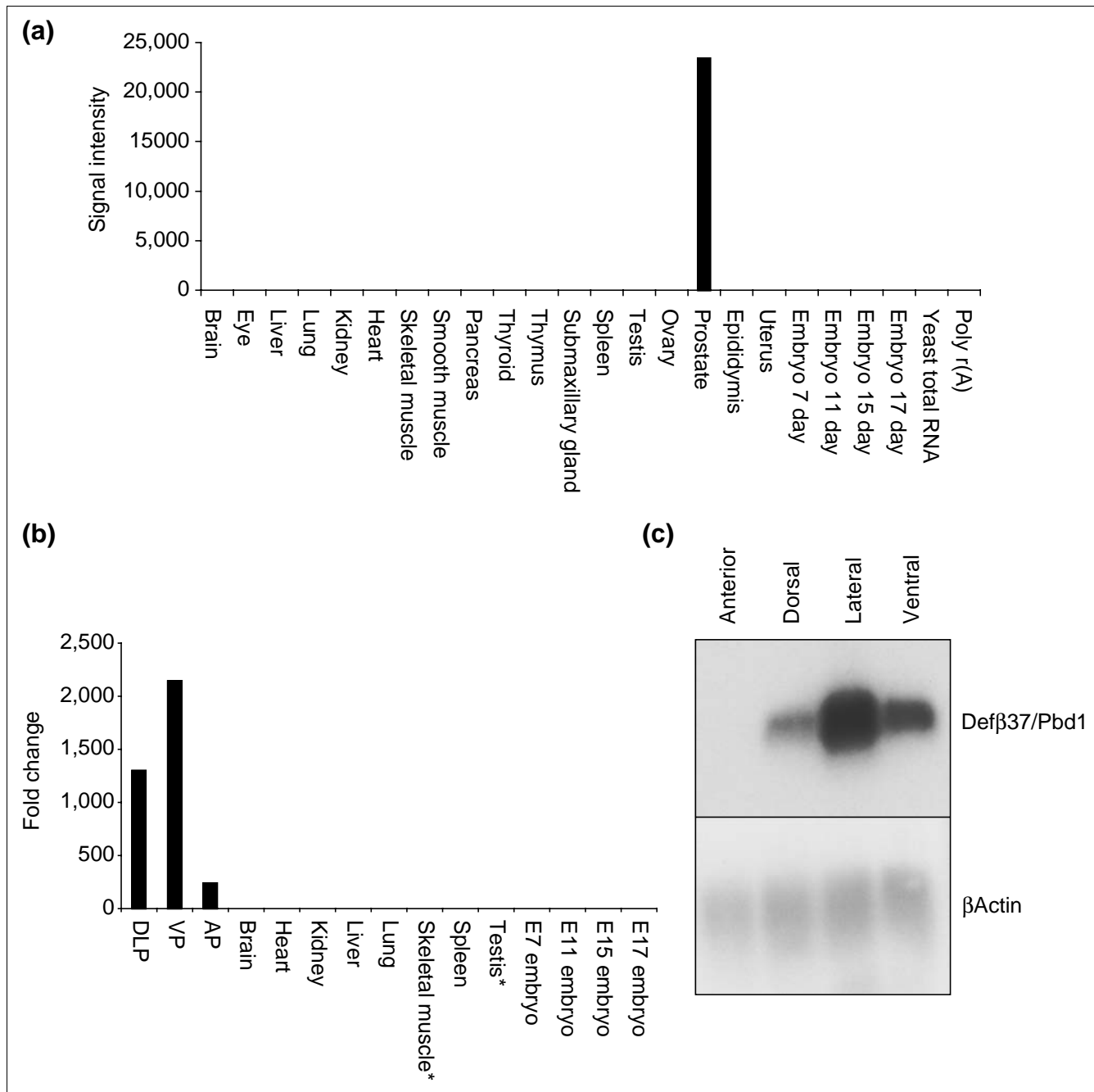
5' UTR and 111 nt 3' UTR with putative polyadenylation sequences located 99, 72 and 51 nt upstream of the polyA tail. Protein sequence alignments with members of the mouse and rat  $\beta$ -defensin family demonstrate a high level of sequence similarity including conservation of the majority of the cysteine and positively-charged residues that are predicted to modulate antimicrobial activity (Figure 6) [35,36]. Alignments of the TU23348 transcript with the draft mouse genome sequence indicates that TU23348 is located on chromosome 8 and is comprised of two exons; the first encoding 19 amino acids and the second 54 amino acids. The exons are separated by a 7.3 kb intron (Figure 6). The gene is located between two other members of the  $\beta$ -defensin family,  $\beta$ -defensin 1 and 2. We named this gene *Pbd1* for prostate  $\beta$ -defensin 1. Subsequently, in accordance with the Human Genome Organization (HUGO) nomenclature committee, this gene carries the official designation of *Defb37* for

defensin beta 37 as it is the 37th  $\beta$ -defensin reported. The sequence is represented in GenBank under accession number AY387658.

### Discussion

The physiological and biochemical features of a particular tissue or cell type represent the complex endpoint of interactions between specific cohorts of expressed genes and the environment. The identification of the complete set of genes expressed in the mouse has been the focal point of intensive efforts involving the sequence analysis of hundreds of cDNA libraries derived from tissues at different stages of mouse development [15]. The ultimate success of this approach depends on the specific organs, tissues and cell types selected for sampling. Many transcripts will be expressed in a tissue- or cell type-restricted manner, while others are expressed





**Figure 5** Prostate-specific expression of a novel  $\beta$ -defensin gene (*Defbeta37/Pbd1*). **(a)** Dot blots comprised of mRNAs derived from multiple normal mouse tissues were hybridized with probes encoding the *Defbeta37/Pbd1* gene, exposed to a phosphorimage screen, and quantitated. Tissue signal intensities are reported in arbitrary units above background intensity. **(b)** Prostate-specific expression of a novel  $\beta$ -defensin gene (*Defbeta37/Pbd1*). QPCR analysis of *Defbeta37/Pbd1* expression in multiple normal mouse tissues and in dissected mouse prostate lobes. Expression levels reflect the relative fold-differences in transcript abundance between tissue samples. Very low levels of *Defbeta37/Pbd1* expression not visible on this plot were detectable in skeletal muscle and testis (asterisks). **(c)** Prostate-specific expression of a novel  $\beta$ -defensin gene (*Defbeta37/Pbd1*). Northern analysis of *Defbeta37/Pbd1* expression in dissected individual mouse prostate lobes.

prostate maturation and the production of testosterone at puberty. These genes include several that encode seminal fluid proteins such as *Svs2*, *Svs5* and *Sbp*. The expression of probasin, a gene known to be regulated by androgens,

followed a similar pattern of expression. Transcripts encoding the antioxidant protein peroxiredoxin 6 (*Prdx6*) (alias antioxidant protein 2: *Aop2*) also increased with age. *Prdx6* is involved in the redox regulation of the cell and can reduce







pairwise comparisons [60], and we report the results of this analysis using a significance level of  $p = 0.001$  with the Bonferroni correction to adjust for multiple comparisons. The DLP01, CG02, VP01 and VP02 libraries were evaluated in a pairwise fashion for the lobular comparison. Libraries included in the developmental pairwise comparison were UGS01, UGS02, NEONATAL, DAY10, DAY20, DAY35, a pooled library for MONTH3 (represented by combined TUs from the lobular comparison) and MONTH14. Genes and ESTs exhibiting differential expression between distinct developmental time points were grouped by hierarchical clustering using the Cluster software [61]. Clusters were visualized using Treeview [61].

Unannotated, putatively novel sequences determined to be differentially expressed between prostate lobes were further evaluated using PSI-BLAST (Position-Specific Iterated BLAST) and PHI-BLAST (Pattern-Hit Initiated BLAST) [58]. Nucleotide sequences were translated using the ExpASY Translate program [62] to identify open reading frames. ESTs were also compared against the assembled mouse genome (February 2003 freeze) through the University of Santa Cruz's BLAST-Like Alignment Tool (BLAT) [56] to identify spliced sequences and predicted genes.

#### Quantitative polymerase chain reaction

Quantitative PCR analysis of selected genes was performed to confirm the virtual expression results. Approximately 2 ng of cDNA from each prostate lobe was used as a template for PCR reactions using buffers and instructions in accordance with the manufacturer's protocol (Applied Biosystems Inc., CA, USA). Reactions contained 1X SYBR Green master mix (Applied Biosystems, Inc.), and 0.3  $\mu$ M of oligonucleotide primer pairs designed to amplify the following genes: probasin (*Pbsn*): Forward- 5' GGTCATCATCTCCTGCTCA 3', Reverse- 5' AGGCCGTCATCTTCTTTTT 3' (79 bp amplicon); spermine binding protein (*Sbp*): Forward- 5' CCCA-CATGCAGAGCCAGAAA 3', Reverse- 5' ATCCGCATGCCCTTGAGTTG 3' (95 bp amplicon); serine protease inhibitor, Kazal type 3 (*Spink3*): Forward- 5' TAT-AGTTCTTCTGGCTTTTGC 3', Reverse- 5' TCTATGCGTTTC-CTGTTTTCA 3' (246 bp amplicon); onzin (alias placenta-specific 8 (*Plac8*)): Forward- 5' TTCTGTCCTGTTTGCTCT-GTG 3', Reverse- 5' TCATGGCTCTCCTCTGTTA 3' (61 bp amplicon); receptor (calcitonin) activity modifying protein 2 (*Ramp2*): Forward- 5' CTCATCCTTCCCACAGACCT 3', Reverse- 5' TGTGTCGTGAGTCCCCTTTG 3' (61 bp amplicon); ventral prostate predominant 1 (*Vpp1*): Forward- 5' TGCTGTCTGTCTGCTTCTG 3', Reverse-5' CCATACTTATT-GTTTCTCCTTTC 3' (126 bp amplicon); ribonuclease, RNase A Family, 1 (*RNase1*): Forward- 5' AAGTCCCTCATTCTGTT-TCCA 3', Reverse- 5' TATCCCGCGTTTCATCATTT 3' (166 bp amplicon); EST similar to acrosomal vesicle protein 1: Forward- 5' TGTTCCCTAGGCTCTCACTGC 3', Reverse- 5' CCAAGAGTAGCAACAAGAGG 3' (52 bp amplicon); prostate  $\beta$ -defensin 1 (*Pbd1*): Forward- 5' GATCAAGTGTAT-

GCCAAAATG 3', Reverse- 5' TTTATATGGCTTCAGT-GCTCTA 3' (149 bp amplicon). Amplifications were performed using a protocol of 95°C for 30 sec, 60°C for 30 sec, and 72°C for 30 sec for 40 cycles on the ABI 7700 sequence detector while recording fluorescence measurements. Each reaction was performed in triplicate and cycle numbers were normalized to a parallel ribosomal protein S16 control as previously described [63]. To determine the tissue distribution of *Pbd1*, quantitative PCR was performed using cDNAs derived from multiple mouse tissues according to the supplier's protocol (Invitrogen).

#### Northern and dot blot analysis

Ten  $\mu$ g of total RNA from each prostate lobe and other mouse tissues were fractionated on 1.2% agarose denaturing gels and transferred to nylon membranes by a capillary method [64]. The mouse multiple tissue and master blots were obtained from Clontech, CA, USA. Blots were hybridized with *Def $\beta$ 37/Pdb1* cDNA probes labeled with [ $\alpha$ -<sup>32</sup>P]dCTP by random priming using the Rediprime II random primer labeling system (Amersham, CA, USA) according to the manufacturer's protocol. Blots were stripped and re-probed for  $\beta$ -actin as a loading control. Filters were imaged and quantitated by using a phosphor-capture screen and ImageQuant software (Amersham, CA, USA).

#### Additional data files

The additional data files are available with this article and at [17]: A table showing the genes that are differentially expressed between developmental timepoints at a significance level of  $p = 0.001$ , without the correction for multiple comparisons (Additional data file 1), and a table of genes represented by more than two prostate ESTs and expressed in fewer than two tissues in addition to the prostate (Additional data file 2).

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