Temporal blastemal cell gene expression analysis in the kidney reveals new Wnt and related signaling pathway genes to be essential for Wilms’ tumor onset

M Maschietto1, AP Trape1,2, FS Piccoli1, TI Ricca1, AAM Dias3, RA Coudry4, PA Galante5, C Torres5, L Fahhan7, S Lourenço4, PE Grundy8, B de Camargo9, S de Souza9, EJ Neves7, FA Soares4, H Brentani10 and DM Carraro*,1

Wilms’ tumors (WTs) originate from metanephric blastema cells that are unable to complete differentiation, resulting in triphasic tumors composed of epithelial, stromal and blastemal cells, with the latter harboring molecular characteristics similar to those of the earliest kidney development stages. Precise regulation of Wnt and related signaling pathways has been shown to be crucial for correct kidney differentiation. In this study, the gene expression profile of Wnt and related pathways was assessed in laser-microdissected blastemal cells in WTs and differentiated kidneys, in human and in four temporal kidney differentiation stages (i.e. E15.5, E17.5, P1.5 and P7.5) in mice, using an orthologous cDNA microarray platform. A signaling pathway-based gene signature was shared between cells of WT and of earliest kidney differentiation stages, revealing genes involved in the interruption of blastemal cell differentiation in WT. Reverse transcription-quantitative PCR showed high robustness of the microarray data demonstrating 75 and 56% agreement in the initial and independent sample sets, respectively. The protein expression of CRABP2, IGF2, GRK7, TESK1, HDGF, WNT5B, FZD2 and TIMP3 was characterized in WTs and in a panel of human fetal kidneys displaying remarkable aspects of differentiation, which was recapitulated in the tumor. Taken together, this study reveals new genes candidate for triggering WT onset and for therapeutic treatment targets.

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Conserved mechanisms that underlie the relationship between normal development and tumorigenesis may give insights into the earliest events that drive transformation. Normal kidney differentiation requires a fine regulation of morphogenetic events, and its deregulation may lead to kidney abnormalities, including agenesis, dysplasia, hypoplasia and tumors.1 Wilms’ tumor (WT), or nephroblastoma, arises from metanephric blastematic cells that are unable to complete the mesenchymal–epithelial transition (MET) resulting in a tumor with varying proportions of undifferentiated blastematous and stromal cells and primitive epithelial structures. These histological features resemble fetal kidneys (FKs) and make WT a model for morphological recapitulation of the earliest kidney developmental stages.1,2

Additional evidence for recapitulation of the earliest kidney differentiation stages in WT has been provided by molecular studies, exemplified by the WT1, IGF2 and other genes, implicated in nephrogenesis and associated with Wilms’ tumorigenesis.2-3 A high degree of similarity has also been detected in gene expression profiles of the earliest kidney differentiation stages and whole WT.4,5 More recently, we have provided evidence that, among the three histological WT components, blastemal cells display a gene expression profile similar to that found in the initial stages of nephrogenesis, suggesting that the blastemal component, and not the whole tumor, preserves the molecular events involved in WT onset.6

Kidney differentiation involves complex interactions between epithelial and mesenchymal cells, and signal transduction pathways have a fundamental role in controlling cell–cell interactions. The Wnt signaling pathway, an inter-related pathway with key roles in embryogenesis and cancer, consists of two highly connected main branches, namely the CTNNB1-dependent and the CTNNB1-independent WT pathways. The CTNNB1-dependent pathway is essential for controlling embryonic development and adult cell homeostasis. β-Catenin, encoded by CTNNB1, functions as coactivator of the TCF/LEF family of transcription factors during embryonic development and is a key component of cell–cell

1Laboratory of Genomics and Molecular Biology, CIPE—AC Camargo Hospital, São Paulo, SP, Brasil; 2Departamento de Radiologia e Oncologia—Faculdade de Medicina da Universidade de São Paulo (FMUSP), São Paulo, SP, Brasil; 3Laboratory of Experimental Genetics, Department of General Biology—Universidade Federal de Minas Gerais, Belo Horizonte, MG, Brasil; 4Department of Anatomic Pathology—AC Camargo Hospital, São Paulo, SP, Brasil; 5Laboratory of Bioinformatics—Ludwig Institute for Cancer Research, São Paulo, SP, Brasil; 6Laboratory of Bioinformatics, CIPE—AC Camargo Hospital, São Paulo, SP, Brasil; 7Instituto de Matemática e Estatística da Universidade de São Paulo (IME-USP), São Paulo, SP, Brasil; 8Pediatric Hematology, Oncology and Palliative Care—Stollery Children’s Hospital, Edmonton, Alberta, Canada; 9Research Pediatric Hematology-Oncology Program, Instituto Nacional do Cáncer, Rio de Janeiro, RJ, Brasil and 10Departamento de Psiquiatria—Faculdade de Medicina da Universidade de São Paulo (FMUSP), São Paulo, SP, Brasil. *Corresponding author: DM Carraro, Laboratory of Genomics and Molecular Biology, Rua Prof. Antonio Prudente, 109—1 andar, CIPE—Hospital A.C. Camargo, CEP 01509-010 São Paulo, SP, Brasil. Tel: +55 11 2189 5023; Fax: +55 11 2189 5185; E-mail: dirco.carraro@cipe.acacamargo.org.br

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Abbreviations: ACIS, Automated Cellular Imaging System III; aRNA, amplified RNA; DK, differentiated kidney; FFPE, formalin-fixed, paraffin-embedded; FK, fetal kidney; hcrHMs, conserved regions of human and mouse orthologs; HM, human and mouse; MET, mesenchymal–epithelial transition; RT-qPCR, reverse transcription-quantitative PCR; TMA, tissue microarray; WT, Wilms’ tumor

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adhesion complexes in differentiated cells. Wnt ligands activate the CTNNB1-dependent pathway, leading to nuclear accumulation of β-catenin, which acts as a coactivator for transcription factors. In the absence of Wnt ligands, this pathway is inactivated by a destruction complex that phosphorylates β-catenin, leading to ubiquitination, proteasomal degradation and prevention of β-catenin accumulation in the nucleus.7 Therefore, nuclear accumulation of β-catenin is an indirect marker for pathway activation and has been observed in WT.8–10 APC, a component of the β-catenin destruction complex, is also found in the nucleus of blastemal cells in the earliest embryonic kidney stages and in WT.6 The CTNNB1-independent Wnt pathway comprises two segments, namely the planar cell polarity and calcium signaling segments,11 and its independent Wnt pathway comprises two segments, namely and prevention of β-catenin accumulation in the nucleus. Therefore, nuclear accumulation of β-catenin is an indirect marker for pathway activation and has been observed in WT.8–10 APC, a component of the β-catenin destruction complex, is also found in the nucleus of blastemal cells in the earliest embryonic kidney stages and in WT.6 The CTNNB1-independent Wnt pathway comprises two segments, namely the planar cell polarity and calcium signaling segments,11 and its proper regulation has also been considered important in maintaining correct cell differentiation in the kidney. The calcium signaling components PLCγ29 and WNT5A12 are downregulated in WT, and their fine regulation seems to have an important role in kidney differentiation.

The strong connection between nephrogenesis and WT highlights the importance of characterizing gene expression modulation in the Wnt and related signaling pathways. Identification of genes expressed during kidney differentiation arrest may be necessary to determine the earliest alterations that drive WT onset and, consequently, are potential molecular markers or therapeutic target candidates. In this study, we characterized gene expression patterns of the Wnt and related signaling pathways in blastemal cells in kidney developmental stages in mice and in WT in humans. For this purpose, we established a cross-species hybridization model to assess the gene expression modulation in both species on a cDNA platform. Our findings revealed 18 genes commonly regulated in the earliest kidney developmental stages and in WT. Expression patterns based on this gene set could discriminate blastemal WT cells and FK cells from those of differentiated kidneys (DKs) from both species. Detailed characterization of the protein expression of human FKs and WT revealed some interesting aspects of gene modulation in nephrogenesis and in WT. Overall, we revealed new genes that determine kidney differentiation arrest and are potential molecular markers or therapeutic target for WT patients.

Results

Evaluation of cross-species hybridization for the construction of a cDNA platform. To test the robustness of interspecific hybridizations, we interrogated 1334 sequences classified as highly conserved regions of human and mouse orthologous using RNA from kidney and liver of both species (see Material and Methods). Correlation between technical replicates and variance among the triplicate hybridization experiments were compared between interspecific (mouse RNA samples and human cDNA platform) and intraspecific (human RNA samples and human cDNA platform) hybridizations, with the latter considered the gold-standard experiment.

Technical replicates of interspecific hybridizations generated highly reproducible data, with Pearson’s correlations ranging from 83 to 86% (average = 84.7%), similarly to intraspecific hybridizations, which resulted in 86–95% (average = 91.3%). To evaluate variance across the experiments, Levene’s test was used to compare the fluorescence intensities of each spot in the intraspecific and interspecific hybridization replicates. For 90.6% of the spots (1195 out of 1334), the variability across the interspecific hybridizations was equal to or lower than that across the intraspecific hybridizations.

These analyses indicated that the criteria for defining highly similar orthologous sequences and the hybridization conditions resulted in robust interspecific hybridization data that were similar to data obtained for the intraspecific hybridizations.

Therefore, the same criteria for defining similar regions in the cDNA sequences (i.e. an e-value < 10−5 and a minimum alignment of 50%) were used for constructing the orthologous cDNA platform used in this study. In total, 326 orthologous cDNA partial fragments with high similarity at the nucleotide level between human and mouse (HM) sequences were immobilized on a custom cDNA platform (Supplementary File 1) to assess gene expression regulation in both species.

Characterization of differentially expressed genes between WT and DK. From the 326 Wnt and related signaling pathway genes assessed in WT and DK, 47 were differentially expressed; and 18 and 29 were downregulated and upregulated in WT, respectively (P<0.05; Table 1).

Annotation of the biological processes classified the 47 differentially expressed genes into cell communication, development, organogenesis, morphogenesis, phosphorylation, regulation of transcription, transport and other categories without any detectable preferential distribution (Table 2).

Identification of genes involved in the earliest event that drives WT onset. To identify genes that recapitulate the expression profile of the earliest kidney developmental

| Downregulated genes in WT | Upregulated genes in WT |
|--------------------------|-----------------------|
| CDC7                     | ITPR3                 |
| CDH6                     | MAPK9                 |
| CDS1                     | OCRL                  |
| DVL2                     | PIK3CA                |
| DVL2                     | PIK3CD                |
| EYA4                     | PIP5K2B               |
| FN3K                     | PIP5K2C               |
| FRAT2                    | TBL1XR1               |
| HIPK1                    | TIMP3                 |
| VANGL2                   | WNT3                  |
| WNT5B                    |                       |
| COL5A1                   | INPP5B                |
| EYA3                     | LRP6                  |
| EYA4                     | PIP5K2B               |
| FN3K                     | PAX2                  |
| FRAT2                    | PCTK2                 |
| HIPK1                    | FZD10                 |
| CARKL                    | DMPK                  |
| INPP5B                   | CRABP2                |
| LRP6                     | FRZ2B                 |
| PIP5K2B                  | GRK7                  |
| PAX2                     | IGFI1R                |
| PCTK2                    | LRP1                  |
| FZD10                    | FZD4                  |
| CSNK2A1                  | OFD1                  |
| HDGF                     | PPP2R1A               |
| JUN                      | TCF7L2                |
| KIAF2                    |                       |
stages and consequently drive tumorigenesis in WT, genes downregulated and upregulated in WT, compared with DK, were compared with genes with increasing and decreasing behavior during kidney cell differentiation. The 326 genes were evaluated in 4 stages of mouse nephrogenesis, which represent the differentiation of metanephric blastema into fully DK cells, revealing 72 and 64 genes with increasing and decreasing expressions, respectively (Supplementary Figure 1). The others 190 genes presented diverse expression behavior during kidney cell differentiation and were not further analyzed.

Thus, the 72 increasing genes were compared with the 18 WT blastemal cell downregulated genes, and the 64 decreasing genes were compared with the 29 WT blastemal cell upregulated genes. This analysis revealed that 18 genes, 7 with increasing and 11 with decreasing expressions during mouse metanephric blastema cell differentiation (Figure 1a), were downregulated (TIMP3, HIPK1, CDH6, ITPR3, FRAT2, PIK3CA and MAPK9) and upregulated (PAX2, FZD10, CARKL, WNT5B, FZD2, CRABP2, FR2B, GRK7, TESK1, HDGF and IGF2) in WT blastemal compared with DK cells, respectively.

Hierarchical clustering based on the expression pattern of the 18 genes grouped differentiated and intermediate kidneys from humans and mice together and discriminated them from WT samples, which were grouped with the pool of human FKs and the earliest mouse embryonic kidneys (Figure 1b). This result revealed two important aspects of this study: the model for interspecific hybridization was appropriate for studying human tumorigenesis and mouse embryogenesis, and a gene expression signature linked to Wnt and related signaling pathways was associated with WT onset.

A high level of robustness was detected in this study, with 75% agreement between microarray and reverse transcription-quantitative PCR (RT-qPCR) experiments in the initial sample set. In total, 9 out of 12 genes displayed concordant expression. CRABP2, HDGF, IGF2, PAX2, SHPK, TESK1 and WNT5B were upregulated and HIPK1 and TIMP3 were downregulated, in WT blastemal cells compared to DK glomerular cells (Figure 1c). Interestingly, hierarchical clustering based on the 12 genes assessed by RT-qPCR, independent of their validation, could discriminate DK from 100% of WT samples (Supplementary Figure 3) reinforcing the association of this gene signature with WT.

### Evaluation of expression levels using a broader group of WT and DK samples.

To assess the transcriptional difference of the 9 technically validated genes (CRABP2, IGF2, PAX2, WNT5B, TIMP3, HIPK1, TESK1, SHPK and HDGF) in samples with different biological backgrounds, we evaluated an independent group of 35 blastema-enriched WT and 13 cortex-enriched DK samples. The data from CRABP2, HDGF, IGF2, TIMP3 and WNT5B agreed with the initial sample group. HIPK1 did not satisfy the fold-change criteria ($\geq 3$), despite showing statistical significance (Figure 2).

### Characterization of protein expression in WT and during human kidney embryogenesis.

First, correspondence between the mRNA and protein was evaluated based on the immunohistochemical labeling intensity of each protein in the blastemal WT and glomerular DK cells. Proteins encoded by 8 out of 18 genes were evaluated in 145 WT and 20 DKs. A high percentage of agreement was observed, with 5 out of 8 (62.5%) genes displaying the same expression tendency at mRNA and protein levels. CRABP2, HDGF, IGF2, TIMP3 and WNT5B had overexpression in WT. TIMP3 and FZD2 did not show a difference between WT and DK, and TIMP3 had opposite results, that is, its mRNA was underexpressed in WT, and its protein product was overexpressed (Figure 3).

Second, protein positivity was characterized in embryonic kidneys, from undifferentiated mesenchymal metanephric to fully differentiated cells. In addition, WT and DK samples were classified as either negative or positive for each protein (Table 3).

A protein expression profile that recapitulated the earliest kidney developmental stages could be observed in WT for the majority of the eight assessed proteins (Figure 3).

Cytoplasmic and some nuclear CRABP2 positivity was detected in every cells undergoing MET after ureteric bud induction and before cells adopted an epithelial morphology. CRABP2 was not detected at any of the subsequent cell differentiation stages. The majority of WT samples displayed strong CRABP2 expression in the nucleus (107 out of 113 samples) and in the cytoplasm (101 out of 108 samples).

With regard to IGF2, cytoplasmic positivity was observed in all embryonic kidneys, with a slightly stronger positivity in the earliest stages. DK cells were considered IGF2 negative, and in WTs, IGF2 expression was observed in 96 out of 112 blastemal cells.

GRK7 was observed in the cytoplasm and in membranes of all embryonic kidneys, showing a slight decrease in the DK. In DK cells, half of the glomerular cells reported positive cytoplasm labeling, and the majority of cells also presented membrane positivity. All but one blastemal WT sample were positive for GRK7 in the cytoplasm (105 out of 106 samples) and in the membrane (116 out of 117 samples).

### Table 2: Classification of differentially expressed genes between relapse and no-relapse WT samples within the biological process categories

| Biological process categories | Upregulated genes in WT | Downregulated genes in WT |
|--------------------------------|-------------------------|---------------------------|
| Cell communication | INPP5B, PIP5K2B, RGS4 | CDH6, CDS1, DYL2, FRAT2, PIK3CA, WNT3, WNT5B |
| Development, organogenesis, morphogenesis | EYA3, FRZB, FZD10, FZD2, FZD4, GRK7, IGF2, PAX2, PPP2R1A, SFRP2, WNT5B | EYA4, VANGL2 |
| Phosphorylation | CSNK2A1, DMPK, IGF1R, PCTK2, PPP2R1A, TESK1 | HIPK1, MAPK9, PIP5K2C, PIK3CD |
| Regulation of transcription | COL5A1, HDGF, JUN, TCF7L2 | ITPR3 |
| Transport | CRABP2, LRP1, PIK3C3 | CDC7, FN3K, OCRL, TBL1XR1, TIMP3 |
| Others | CARKL, LRP6, OFD1 | |
**Figure 1** Wilms tumor signature set. (a) Expression behavior of the 18 genes during nephrogenesis in mouse. ▼ and ▲ indicate downregulation and upregulation in Wilms tumor, respectively, in comparison with differentiated kidney at transcriptional level. (b) Unsupervised hierarchical clustering based on expression pattern of the 18 genes. Black lines correspond to 90–100% of reliability assessed by Bootstrap. (c) Comparison of cDNA microarray and RT-qPCR experiments. (*) Represents differentially expressed genes assessed by RT-qPCR that fulfilled cutoff criteria (fold change > 3 and P < 0.05). (d) Legend of colors. FK, fetal kidney; WT, Wilms tumor; DK, differentiated kidney; P1.5, 1.5-day-old mouse kidney; E17.5 and E15.5, fetal kidney from 17.5 and 15.5-day-old mouse embryo, respectively.
TESK1 exhibited a weak cytoplasmic positivity, primarily in tubular cells and a few embryonic kidney glomerular cells; in DKs, some glomerular cells were also positive. TESK1 showed positive staining in the cytoplasm of the majority of the WT blastemal components (110 out of 114 samples).

HDGF nuclear positivity was observed in some cells from all FK stages and DK. HDGF was positive in the cytoplasm of 56 out of 118 cells and in the nucleus of 52 out of 111 WT blastemal components.

TIMP3 was labeled in cell membranes and in the nuclei of embryonic kidney cells. TIMP3 was also observed in the cytoplasm and nucleus of DKs and in WT blastemal cells; however, stronger labeling was observed in the latter. Whereas the majority of samples were positive in the cytoplasm (101 out of 116 samples) less than half were positive in the nucleus (47 out of 111 samples).

In nephrogenesis, WNT5B was observed after renal vesicle formation, when cells are organizing to form kidney structures, such as glomeruli and tubules. WNT5B positivity was observed mainly in the apical cell membrane. In DK, WNT5B nuclear positivity was observed in some cells. In the WT blastemal component, WNT5B was observed in the cytoplasm in 8 out of 120 samples and in the membrane in 28 out of 113 samples.

With regard to FZD2, FKs were membrane positive in cell types ranging from undifferentiated metanephric blastema cells to differentiated glomerular and tubular cells. In WT blastemal cells, FZD2 was observed in the membrane in 37 out of 106 samples.
Assessment in relapse and no-relapse WT samples. To determine whether the 9 technically validated genes (i.e. CRABP2, IGF2, PAX2, WNT5B, TIMP3, HIPK1, TESK1, SHPK and HDGF) had a role in the latter events of WT progression, their expression levels were assessed in 35 blastema-enriched WT samples that included 12 relapse and 23 no-relapse samples (Supplementary Figure 4). Only IGF2 was differentially expressed ($P<0.01$); presenting downregulation in relapse WTs. TESK1 was also downregulated in relapse WTs, but the $P$-value was borderline ($P=0.06$).

In addition, in an effort to find prognostic markers for WT, possible associations between protein expression (CRABP2, GRK7, HDGF, IGF2, TESK1, WNT5B, FZD2 and TIMP3) and clinicopathological variables (relapse, clinical stage, first treatment, number of drugs used, death, age of diagnosis and gender) were evaluated. No association was found, reinforcing that these molecular events had a role rather in tumorigenesis than in progression of WT.

**Discussion**

Embryonic tumors are intimately linked to embryogenesis, tissue growth and maturation process. Embryogenesis and tumorigenesis have been shown to share regulatory mechanisms in several tumors including WT.\(^4\)\(^6\) Approaches that enable the assessment of both processes have been routinely used to uncover key molecular events that lead to cell transformation.

Nephrogenesis is a continuous process that involves different cell types, in variable proportions, the spatiotemporal localization of which has a role in the correct differentiation of the functional kidney. The triphasic WT histology resembles FK cells.\(^13\) We previously analyzed the gene expression pattern of blastemal, stromal and epithelial components of WT separately and showed that blastema displays the most similar gene expression pattern of the initial stages of nephrogenesis, and that the Wnt signaling pathway is supposedly involved in the earliest events of WT onset.\(^6\)

In the current study, a cell-resolution expression analysis was used for identifying disturbances in gene modulation of blastemal cell differentiation that can trigger a permissive condition for tumor development. We analyzed the expression modulation of Wnt and related signaling pathways genes in primordial blastemal WT cells and in four FK stages to identify conserved mechanisms in the earliest stages of kidney embryogenesis that recapitulate Wilms’ tumorigenesis. The rationale behind this experimental design was to define alterations in gene expression modulation that are involved in the disruption of the differentiation process in the kidneys that lead to WT onset. To monitor gene expression in both processes, that is, tumorigenesis in humans and embryogenesis in mice, we used stringent parameters and generated reliable data from cross-species hybridization.

The signature of 18 Wnt and related signaling pathways genes displayed an expression pattern that was modulated during the differentiation of blastemal cells in nephrogenesis. Additionally the expression pattern of blastemal cells from the earliest stages of nephrogenesis in mice was recapitulated in blastemal cells from the WT in human. These results reinforced the importance of these genes in WT onset and also validated the approach used in this study.

Activation of the canonical Wnt signaling pathway has been continuously detected in WT verified by nuclear positivity of APC\(^6\) and $\beta$-catenin.\(^9\)\(^\text{—}\)\(^14\) In spite of the fact that no experimental evidence has been provided in this study to connect the 18 genes and activation of the canonical Wnt pathway, it is reasonable to suggest their involvement with this process in both nephrogenesis and WT.

Among the nine technically validated genes, PAX2, IGF2, CRABP2, HDGF, SHPK, TESK1 and WNT5B were upregulated and HIPK1 and TIMP3 were downregulated at the mRNA level in WT blastemal cells than in DK glomerular cells. PAX2 (located at 10q24) is a transcription factor that is expressed in FK nephrogenic structures and in WT, but is absent from the DK.\(^15\) PAX2 expression is suppressed by WT1, and both are essential for kidney development.\(^16\) Inactivating WT1 mutations have been observed in $\sim$ 10% of sporadic WT cases,\(^7\) and germline mutations are associated with syndromes that include an increased WT incidence.\(^13\) Consistent with these data, PAX2 was similarly upregulated in this study at the mRNA level in WT as compared with the earliest FK developmental stages.

IGF2 (located at 11p15.5) is an embryonic growth factor and a member of the insulin family of polypeptides, with a role in kidney embryogenesis.\(^18\) Under physiological conditions, the IGF2 maternal allele is silenced by genomic imprinting, and the gene is expressed only by the paternal allele; however, WT exhibits a markedly increased expression relative to DK.\(^19\)\(^20\) As expected, IGF2 upregulation was detected in the earliest nephrogenesis stage, and this was recapitated by WT both at the mRNA and at the protein levels.

WNT5B (located at 12p13.3) encodes a secreted Wnt signaling protein. CGH arrays of WTs have revealed a high level of gain of the WNT5B chromosomal locus,\(^21\) which is in accordance with the transcriptional upregulation found in our study. However, no alteration at the protein level was observed in our results.

In addition, amplification of the chromosome region 1q22-q23.1, where HDGF and CRABP2 are located, was also observed in WT samples,\(^21\) and this is in agreement with the

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**Figure 3** Representative sections of immunohistochemical staining in Wilms tumor, fetal and differentiated kidneys. Proteins are identified in each block. Antibody specificity was determined by western blot (panels above graphics). Cell line and molecular weight are indicated. HEK293: human embryonic kidney cell line, SK-NEP-1: Wilms tumor cell line. Graphics: Y axis—protein quantification based on protein intensity values. X axis: Wilms tumor (right) and differentiated kidney (left) samples. A, D, G, J, M, P, S and V: representative samples of Wilms tumor. B, E, H, K, N, Q, T and X: representative samples of fetal kidneys. Fetal kidneys from thirteenth week human embryo: HDGF, IGF2, GRK7, WNT5B and TIMP3; fetal kidneys from seventeenth week human embryo: CRABP2 and FZD2; and; fetal kidneys from the twenty-fourth week human embryo: TESK1, C, I, L, O, R, U, Y: representative samples of differentiated kidney, only cells from glomerulocystic were analyzed. WT: Wilms tumor; DK: differentiated kidney.

(*) Represents differentially expressed genes with $P<0.05$. Black bars represent 100.1 or 100.3 $\mu$m as indicated in the green box.
Figure 3  Continued

WT and kidney development
M Maschietto et al

HDGF
SK-NEP-1  37kDa

WNT5B
HEK283  40kDa

FZD2
HEK293  63kDa

TIMP3
HEK293  33kDa

m
n
o
p
q
r
s
t
u
v
w
x

*p<0.01
P<0.01
P=0.44
P=0.74
P<0.01
overexpression observed in our study. Our data also demonstrate slight HDGF overexpression at the protein level in blastemal cells than in DK glomerular cells, which reinforces the idea that HDFG is downregulated throughout organ differentiation.

In this study, CRABP2 was upregulated both at the mRNA and at the protein level in WT blastemal cells than in cells from the cortex of DK samples in accordance with other studies in WT. However, CRABP2 downregulation has been observed in others tumor types including head and neck squamous cell carcinoma, in which a highly frequent hypermethylation rate was detected. CRABP2 encodes a cytosolic member of the retinoic acid-binding protein family involved with nephron formation through Wnt4 in mouse. In addition, WNT5B belongs to the Wnt calcium signaling pathway, which has been recently shown to be involved in the earliest kidney stages. Indeed, WNT5B is expressed in blastemal cell differentiation, which may be perturbed in WT.

Table 3 Characterization of proteins expression in blastemal cells of WT, glomerulous cells of DK and in different stages of fetal kidneys

| Protein | Localization | WT Negative | WT Positive | DK Negative | DK Positive | Nephrogenesis |
|---------|--------------|-------------|-------------|-------------|-------------|--------------|
| CRABP2  | Cytoplasm    | 6 (5.3%)    | 107 (94.6%) | 18 (100%)   | 0 (0%)      | Cytoplasm expression in cells going through MET |
|         | Nucleus      | 7 (6.4%)    | 101 (93.5%) | 17 (100%)   | 0 (0%)      |              |
| HDGF    | Cytoplasm    | 62 (52.5%)  | 56 (47.5%)  | 12 (100%)   | 0 (0%)      | Nuclear and cytoplasm expression in all stages of kidney differentiation |
|         | Nucleus      | 59 (53.15%) | 52 (46.8%)  | 18 (90%)    | 2 (10%)     |              |
| GRK7    | Cytoplasm    | 1 (0.94%)   | 105 (99.06%)| 11 (55%)    | 9 (45%)     | Decreasing cytoplasm expression in cells during kidney differentiation |
|         | Membrane     | 1 (0.86%)   | 116 (99.16%)| 2 (9.09%)   | 20 (90.91%) |              |
| TIMP3   | Cytoplasm    | 15 (12.90%) | 101 (87.10%)| 26 (83.87%) | 5 (16.13%)  | Decreasing cytoplasmatic expression in blastemal cells during kidney differentiation |
|         | Membrane     | 64 (57.66%) | 47 (42.34%) | 21 (100%)   | 0 (0%)      |              |
| WNT5B   | Cytoplasm    | 112 (93.33%)| 8 (6.67%)   | 10 (100%)   | 0 (0%)      | Protein observed in the apical membrane of lumen cells that originate kidney structures |
|         | Membrane     | 85 (75.22%) | 28 (24.78%) | 12 (60%)    | 8 (40%)     |              |
| FZD2    | Membrane     | 69 (65.10%) | 37 (34.90%) | 3 (18.75%)  | 13 (81.25%) | Membrane expression in all stages of kidney cell differentiation |
| TESK1   | Cytoplasm    | 4 (3.51%)   | 110 (96.49%)| 20 (100%)   | 0 (0%)      | Decreasing cytoplasmic expression in cells during kidney differentiation |
| IGF2    | Cytoplasm    | 16 (14.29%) | 96 (85.71%) | 10 (100%)   | 0 (0%)      | Decreasing cytoplasmic expression in cells during kidney differentiation |

Abbreviation: MET, mesenchymal–epithelial transition

expression at the mRNA level and increased expression at the protein level in WT, suggesting a posttranscriptional mechanism of protein regulation for TIMP3.

GRK7 (located at 3q24) encodes a member of the guanine nucleotide-binding protein (G protein)-coupled receptor kinase subfamily of the serine threonine protein kinase family. FZD2 (located at 17q21.1) encodes a seven-transmembrane domain protein involved in the signaling of both CTNNB1-dependent and CTNNB1-independent Wnt pathways. FZD2 has been shown to be overexpressed in FKs and stem-like WT xenografts. Accordingly, we observed upregulation at the mRNA level in WT and in the earliest kidney stages.

In addition, WNT5B and FZD2 belong to the Wnt calcium signaling pathway, which has been recently shown to be involved with nephron formation through Wnt4 in mouse. Interestingly, blastemal cells that express WNT5B also express FZD2 in FKs, although not all cells that express FZD2 express WNT5B (data not shown). Furthermore, glomerular DK cells and the epithelial component (the most differentiated WT component) express both proteins (data not shown), which supports the involvement of Wnt calcium signaling in blastemal cell differentiation, which may be perturbed in WT.

In spite of the fact that experimental delineation has favored identification of genes involved in the earliest event of WT onset that not necessarily have a role in WT progression, two genes, namely IGF2 and TESK1, presented mRNA downregulation (with marginal statistical significance), showing a slight association with relapse, which has been constantly investigated for better stratifying WT patients. Further studies are necessary to definitively estimate the value of these genes for prognostic marker to be introduced in WT clinical routine.
It is possible that the utility of these gene expression data, in the context of molecular markers, will be in the application of therapeutic WT intervention because expression changes of these 18 genes seem to be in the earliest events in cell transformation. In addition, these alterations are present in the majority of tumor cells that increase their importance as molecular markers.

In summary, this study provides a clearer outline of the disruption of nephrogenesis and reveals new Wnt and related signaling pathway genes, which are highly likely to have crucial roles in WT biology.

Materials and Methods

Establishment of criteria for highly similar cDNA regions of orthologous genes. To identify highly similar regions between HM orthologous genes, 4608 partial human cDNA sequences immobilized on a custom cDNA platform were BLASTed against mouse transcriptome databases (i.e. MGC, RefSeq and mRNAsHTCs) using an e-value of −5 and a minimum alignment of 50% as parameters. This analysis revealed 1334 human sequences with matched mouse orthologous sequences. Visual inspection revealed an average similarity of 87%, which ranged from 79 to 98%, between sequences. These sequences were classified as highly conserved regions of HM orthologs (hcrHMs).

Reproducibility test of HM interspecies microarray hybridization. To determine the ideal hybridization conditions for HM interspecies hybridization, frozen human samples stored at −140 °C from three normal kidneys and three normal livers were obtained from the A.C. Camargo Hospital Tumor Tissue Biobank (São Paulo, Brazil), and total RNA pools were generated. A kidney and a liver were removed from an adult CD1 mouse, and total RNA was isolated using TRIzol (Invitrogen Life Technology, Carlsbad, CA, USA), according to the manufacturer’s protocol.

A two-round RNA amplification was performed for all HM samples. Labeled cDNA was generated by reverse transcription using 7 μg of amplified RNA (aRNA), random hexamer primers (Invitrogen), Cy3- or Cy5-labeled dCTP (Amersham Biosciences, Piscataway, NJ, USA) and SuperScript II (Invitrogen) according to the manufacturer’s protocol. Dye-swap experiments were performed for intraindividual (i.e. human samples and a human cDNA platform) and interspecific (i.e. mouse samples and a human cDNA platform) hybridizations for dye bias control. Prehybridization and hybridization were performed as described previously. After background subtraction, quantified signals were log transformed, and a nonlinear normalization (Lowess) was applied to correct experimental bias within each array. Dye-swap hybridizations were performed in triplicate for HM samples, and generated data were used for mathematical and statistical analyses. The reproducibility of signal intensities was evaluated for the 1334 hcrHMs by Pearson’s correlation coefficients between replicates and Levine’s tests among triplicates. For the latter, spot signal intensity values from HM triplicates were plotted to calculate the S.D. and signal intensity variance. Variance values from the interspecific hybridizations were used as a standard and compared with those from the interspecific hybridizations.

Construction of a cDNA platform for HM interspecies microarray hybridization. To define the hcrHMs of 326 genes representative of Wnt and related signaling pathways (Supplementary File 1), human FASTA sequences were BLASTed against mouse transcriptome databases (i.e. MGC, RefSeq and mRNAsHTCs) using described parameters (>50% identity and an e-value of −5). Primers were designed to flank the hcrHMs of 101 genes, and RT-PCR was performed using cdNAs converted from a pool of total RNA from 15 human cell lines as described previously. The amplicons were extracted from 1% agarose gel and performed using cDNAs converted from a pool of total RNA from 15 human cell lines. A kidney and a liver were removed from an adult CD1 mouse, and total RNA was isolated using TRIzol (Invitrogen Life Technology, Carlsbad, CA, USA), according to the manufacturer’s protocol.

Two pathologists (FAS and RC) supervised all dissection procedures. This study was in accordance with the principles of the Declaration of Helsinki and was approved by the A.C. Camargo Hospital Ethics Committee under number CEP 764/06 for human samples. All animal experiments were also conducted according to the guidelines of the A.C. Camargo Hospital Animal Ethics Committee under the number CEUA 006/06.

RNA extraction and amplification. RNA from laser-microdissected samples was purified using the RNeasy-Free DNase set (Qiagen, Germantown, MD, USA). Total RNA from scalpel-dissected samples was extracted using TRIzol (Invitrogen). To generate aRNA, we used the RiboAmp RNA amplification kit, and the RiboAmp HS plus kit (Arcturus Engineering) for HM samples, respectively. RNA quantity and purity were determined by spectrophotometry NanoDrop ND-1000 (Thermo Scientific, Wilmington, DE, USA). Total RNA integrity was analyzed by Agilent Bioanalyzer 2100 (Agilent Technologies, Palo Alto, CA, USA) and only those with an OD 260/280 ≥ 1.8 and an RIN > 5.0 were used for microarray and RT-qPCR experiments.

cDNA microarray experiments. A pool of RNAs was used as the reference sample. Labeled cDNA was generated using 7 and 4 μg of aRNA from the test and reference samples, respectively, random hexamer primers (Invitrogen), Cy3- or Cy5-labeled dCTP (Amersham Biosciences) and SuperScript III (Invitrogen) for competitive hybridization to the cDNA platform. Dye swapping was performed for each sample to control for dye bias, and this was used as a replicate.

For mouse samples, dye-swap experiments were performed in triplicate for animals aged E17.5, P1.5 and P7.5 and for one replicate of E15.5. Data acquisition and normalization were performed as described previously. The microarray data files were deposited in the GEO/NCBI public database identified as GSE25965 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE25965).

RT-qPCR. To validate the differentially expressed genes from microarray experiments, the initial sample set was evaluated using RT-qPCR, which is referred to here as technical validation. cDNA was synthesized from 2 μg of aRNA using SuperScript III (Invitrogen) and random hexamer primers. For an independent sample set, which is referred to here as biological validation, cDNA was synthesized from 2 μg of DNase-treated total RNA using iRNAPROM II (Promega, Madison, WI, USA) and oligo-dT(24) primers, according to the manufacturer’s protocol. RT-qPCR was performed in the ABI Prism 7900 sequence detection system (Applied Biosystems) using SYBR Green PCR Master Mix (Applied Biosystems) in a 20 μl total volume. All reactions were performed in duplicate, and no-template reactions were used as a negative control. The amplification efficiency of each primer pair was calculated using standard curve dilutions (Table 4). ACTB and GAPDH were
selected as the most stable control genes from four endogenous genes tested (GAPDH, ACTB, HPRT1 and BCR) using the geNorm algorithm.56

TMA construction and immunohistochemistry assays. Formalin-fixed, paraffin-embedded (FFPE) WT and kidney specimens from 145 and 20 patients, respectively, were obtained from the Pathology Department of the A.C. Camargo Hospital. For tissue microarray (TMA) construction, hematoxylin and eosin sections were analyzed, and a representative area of the tumor blastemal component and the kidney cortex region was marked on the slide. Selected areas from each sample were arrayed once with a 1.0-mm diameter core that was spaced 0.2 mm apart using a Tissue Microarrayer (Beecher Instruments, Silver Spring, MD, USA).

Additional slides from 19 FFPE postmortem human fetuses at 13 to 24 weeks (i.e. 13, 17, 18, 19, 20, 21, 22 and 24 weeks) of gestation were obtained from the University of São Paulo Medical School in accordance with the authorization of their ethics committee.

Slides were deparaffinized, rehydrated and then subjected to antigen retrieval. The antigen retrieval method details and primary antibody clones are described in Table 5. Sections were incubated in 3% aqueous hydrogen peroxide for 15 min to quench endogenous peroxidase activity and treated with serum-free protein block (Dako, Copenhagen, Denmark) for 20 min at room temperature to suppress nonspecific binding. Immunohistochemical reactions were performed using a polymer technique (Dako). The slides were incubated with the biotinylated secondary antibody from the LSAB polymer technique (Dako). The slides were then counterstained with hematoxylin.

Western blot analysis was performed to check the specificity of antibodies used in IHC assay. HEK293 (CRL-1573) and SK-NEP-1 (HTB-48) cell lines (American Type Culture Collection, ATCC, Manassas, VA, USA) were maintained in a humidified incubator at 37 °C in the presence of 5% CO2. The HEK293 cell line was cultured in DMEM medium (Gibco, Carlsbad, CA, USA) supplemented with 10% FBS and 1% L-glutamine, and SK-NEP-1 cells were maintained in McCoy’s 5A medium (Gibco) supplemented with 15% FBS and 1% L-glutamine. Total protein were extract from subconfluent culture cells with cold RIPA buffer (50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% SDS and 1% Triton X-100 complete, mini protease inhibitor cocktail (Roche)). Western blot analyses were performed as described previously. Proteins were detected using rabbit polyclonal anti-GRK7 (1 : 500), anti-WNT5B (1 : 100), anti-TIMP3 (1 : 2000), anti-FZD2 (1 : 2000), anti-IGF2 (1 : 1000), anti-TESK1 (1 : 500) and mouse monoclonal anti-CRABP2 (1 : 250) antibodies. Signals were detected using ECL HRP-conjugated immunoglobulin G whole antibodies (1 : 2000, GE Healthcare, Little Chalfont, UK).

Protein localization was determined for the nucleus, membrane and cytoplasm. Immunohistochemical expression was quantitatively analyzed using the Automated Cellular Imaging System III (ACIS) (Dako), which detects hue, saturation and luminosity levels and converts these signals into a numerical density measurement that ranges from 0 to 256. Each cell compartment (nucleus, membrane and cytoplasm) evaluated using the ACIS equipment requires an independent slide scanning. Therefore, the number of high-quality images of each protein able to cell recognition and intensity measurement could vary.

For each of the eight proteins analyzed, the pathologist established a limit intensity value by visual evaluation, and this was considered the cutoff value (Table 5). Samples with intensity values lower than the cutoff and with < 10% of cells labeled were considered to be negatively stained. Accordingly, samples with intensity values higher than the cutoff intensity were considered to be positively stained. For DKS, only glomerular cells were used for this analysis because of a high level of nonspecific labeling of tubule cells.

**Mathematical and statistical analysis.** Student’s t-test was used to identify differentially expressed genes between WT and DK in the cDNA microarray experiments. The differentially expressed genes were annotated in the Gene Ontology Database biological process categories and organized in a hierarchical manner through automated EASE gene function analysis.57 To characterize gene expression modulation during the four stages of nephrogenesis, the average signal...
intensity among triplicates of dye-swap microarray experiments was plotted in graphs to create a curve to report the expression tendency. Pearson’s correlation and complete linkage were used for nonsupervised hierarchical clustering, and reliability was assessed by bootstrapping using TMEV version 4.6.

For RT-qPCR experiments, relative fold changes were calculated considering primer efficiency, and genes were considered to be differentially expressed when they reached the previously described criteria of a F-value < 0.05 by the F-test and a fold change ≥ 2 to avoid artifactual results in the relative gene expression data, as recommended previously.36

Membrane, cytoplasm and nucleus density measurements from immunohistochemistry assays were plotted in a graph, and Student’s t-test was applied to find significant differences in labeling between WT blastemal cells and DK glomular cells (P < 0.05).

Conflict of Interest

The authors declare no conflict of interest.

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