Research Paper

Intra-Tumor Genetic Heterogeneity in Wilms Tumor: Clonal Evolution and Clinical Implications

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A B S T R A C T

The evolution of pediatric solid tumors is poorly understood. There is conflicting evidence of intra-tumor genetic homogeneity vs. heterogeneity (ITGH) in a small number of studies in pediatric solid tumors. A number of copy number aberrations (CNA) are proposed as prognostic biomarkers to stratify patients, for example 1q+ in Wilms tumor (WT); current clinical trials use only one sample per tumor to profile this genetic biomarker. We multisampled 20 WT cases and assessed genome-wide allele-specific CNA and loss of heterozygosity, and inferred tumor evolution, using Illumina CytoSNP12v2.1 arrays, a custom analysis pipeline, and the MEDICC algorithm. We found remarkable diversity of ITGH and evolutionary trajectories in WT. 1q+ is heterogeneous in the majority of tumors with this change, with variable evolutionary timing. We estimate that at least three samples per tumor are needed to detect >95% of cases with 1q+. In contrast, somatic 11p15 LOH is uniformly an early event in WT development. We find evidence of two separate tumor origins in unilateral disease with divergent histology, and in bilateral WT. We also show subclonal changes related to differential response to chemotherapy. Rational trial design to include biomarkers in risk stratification requires tumor multisampling and reliable delineation of ITGH and tumor evolution.

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1. Introduction

Intra-tumor genetic heterogeneity (ITGH) has been documented in several adult tumors. Such tumors typically evolve over long periods before diagnosis, with most demonstrating branched evolutionary trajectories (Nowell, 1976; Greaves and Maley, 2012; Gerlinger et al., 2012, 2014; de Bruin et al., 2014). However, the prevalence and relevance of ITGH are poorly understood in pediatric solid tumors: since they carry lower burdens of mutational changes and have evolved for shorter periods of time before diagnosis, they may be expected to show less complex evolutionary histories (Vogelstein et al., 2013).

Although there are relatively few sequence mutations in pediatric malignancies, DNA copy number aberrations (CNA) and rearrangements are often characteristic features of these tumors. Some common CNA have recognized prognostic significance in pediatric tumors. For example, in neuroblastoma, MYCN amplification or subchromosomal genomic gains and losses are used to stratify therapy (Schlieermann et al., 2011). In Wilms tumor (WT), gain of 1q (1q+) is increasingly being proposed as a common prognostic biomarker to select patients for more intensive treatment (Natrajan et al., 2006; Gratias et al., 2013; Segers et al., 2013). However, these studies have relied on a single tumor sample from each case.

Indeed, there has been limited investigation of ITGH in pediatric solid tumors. A recent multisampling study reported genetic homogeneity in multi-sampled embryonal brain tumors (Remke et al., 2015). However, a study of four pediatric small round cell tumors, with two samples from each, reported heterogeneous CNA in three out of four tumors (Mengelbier et al., 2015).

In WT, a large study showed that combined loss of heterozygosity (LOH) of chromosomes 1p and 16q, while rare, was not only associated...
with poorer outcome, but also showed concordance in the vast majority of the 10% of tumors from which two separate samples were assessed (Grundy et al., 2005). In contrast, heterogeneous WTX deletion has been reported in two multi-sampled WTs (Wegert et al., 2009), and heterogeneous activation of MYCN and inactivation of TPS3 have been reported in a case of bilateral WT (Popov et al., 2013; Williams et al., 2015).

Such variable heterogeneity complicates clinical decision making because of a poor understanding of the evolution of pediatric tumors. It also means that most previous studies showing prognostic significance for specific CNA did not take into account potentially significant ITGH. Therefore, here we assess the extent and significance of ITGH in a prospective study of unselected multi-sampled Wilms’ tumors.

2. Methods

2.1. Samples

We obtained multiple samples from WT nephrectomy/nephron-sparing surgery specimens at Great Ormond Street Hospital between May 2011 and June 2013. All patients were enrolled on the SIOP WT 2001 trial (Pritchard-Jones et al., 2015), the current IMPORT study or their parents had consented for additional tissue to be used in research as part of the UK Children’s Cancer and Leukaemia Group tissue bank. The research reported here was approved by a national research ethics committee. Patients received preoperative chemotherapy as per the SIOP WT 2001 trial protocol or according to national clinical guidelines based on this trial. Tumors were classified as previously described (Vujanić et al., 2002). A histological section from each tissue sample was reviewed to determine viable tumor content, and only samples with more than 50% viable tumor (the remainder consisting of necrotic tumor/post-chemotherapy change) were used. DNA was extracted using standard techniques from each tumor sample, and from adjacent non-tumorous kidney where it was available in 19 cases, and from peripheral blood lymphocytes in 3 cases.

2.2. Imaging

In two cases, different regions within the same tumor were identified prospectively as distinct nodules in the same overall tumor mass on T1- and T2-weighted MR imaging, and matched on comparison of pre- and post-chemotherapy images. Assessed diffusion coefficient (ADC) was calculated by one observer (ØEO) as previously described (McDonald et al., 2010).

2.3. Molecular Analyses

Illumina® HumanCytosNP-12 v2·1 microarrays (~300,000 SNP probes) were hybridized with 250 ng DNA per sample according to the manufacturer’s instructions. Methylation-specific multiplex ligation-dependent probe amplification (MS-MLPA) for 11p15 was carried out as previously described (Scott et al., 2008a), using the Salsa MS-MLPA BWS/KSS ME030-C3 probe mix (MRC-Holland), and data visualized in Coffalyser.NET (MRC-Holland).

2.4. Detection of Allele-Specific Copy Number Aberrations and Inference of Tumor Evolution

Log R ratio (log2[observed intensity/reference intensity], LRR) and B-allele frequency (BAF) were calculated using the Illumina® GenomeStudio software for each array using default settings. LRR genomic waves (Diskin et al., 2008) were detected in normal tissue samples and corrected from all arrays. The LRRs from each array were segmented and copy number states were called using the ‘CGHcall’ R package (van de Wiel and van Wieringen, 2007) in Bioconductor (Gentleman et al., 2004). For each case, the boundaries between adjacent regions were compared, smoothed and summarized between samples using the ‘CGHregions’ R package (van de Wiel and van Wieringen, 2007) in Bioconductor (Gentleman et al., 2004). A region was removed if it contained fewer than 100 probes or its probe density was an outlier. The mean tumor-specific mirrored BAF (mBAF) was calculated for each aberrant region and copy number aberrations were rejected if they did not show expected allelic imbalance. Aberrant regions detectable only in the BAF (i.e. copy number neutral LOH) were incorporated into our analysis using a custom pipeline. Allele-specific copy number was interpreted by the phylogetic algorithm MEDICC (Schwarz et al., 2014) to infer clonal evolution of samples in each case. Normal tissue samples were used to root phylogenetic trees. Annotated code of our entire analysis pipeline is available as a GitHub repository at: https://github.com/luslab/multiregion-cnv-phylogenetics.

2.5. Role of the Funding Source

The study sponsors did not participate in study design, in the collection, analysis, and interpretation of data, in the writing of the report, or in the decision to submit the paper for publication.

3. Results

3.1. Intra-Tumor Genetic Heterogeneity in Wilms Tumor Demonstrates Unexpected Diversity

We studied 70 distinct tumor samples from 24 tumors in 20 patients (mean 3.5 samples/case, range 2–6 samples), with matched DNA from non-tumorous kidney and/or peripheral blood leukocytes in 19 cases. Five patients (Cases 9, 10, 16, 17, 20) had bilateral WT, and we obtained samples from both tumors in four of them; in Case 9, the contralateral tumor had been removed prior to the start of our study. Patient characteristics and samples are summarized in Table 1. We applied a custom-built pipeline to determine reproducibly genome-wide allele-specific CNA and LOH events using high-resolution SNP arrays hybridized with genomic DNA from each sample, and automatically compare these events across samples in a tumor. Fig. 1 shows a graphical representation of all CNA and copy number neutral LOH (CNNLOH) events across the 70 tumor samples. We detected most known recurrent WT CNA/LOH, including those associated with poor outcome (Natrajan et al., 2006; Gratias et al., 2013; Segers et al., 2013; Grundy et al., 2005). Surprisingly, 1q+ was heterogeneous in four of seven (57%) multi-sampled tumors with this change (see below). In general, we found remarkable diversity in the extent of intra-tumor CNA and CNNLOH heterogeneity, ranging from cases with unique CNA/CNNLOH events in all or most samples (such as Cases 1 and 15) to tumors exhibiting no CNA/CNNLOH heterogeneity across samples. The latter either lacked somatic CNAs/CNNLOH (Case 12) or showed a consistent pattern of somatic CNAs/CNNLOH across all samples (Cases 2, 3, 6), and the single dominant clone in each tumor showed few (0–4) CNAs/CNNLOH events. The four patients with these homogeneous (and unilateral) tumors were not statistically significantly younger than the other twelve patients with heterogeneous (and unilateral) tumors were not statistically significantly younger than the other twelve patients with heterogeneous unilateral tumors (Welch two-sample one-tailed t-test, t = −1.52, p = 0.08), suggesting that heterogeneity does not arise purely as a consequence of later age at diagnosis.

3.2. Diverse Evolutionary Pathways in Wilms Tumor: From Homogeneity to Branched Evolution

In order to infer the evolutionary history of tumors, we generated phylogetic trees depicting the relationships between samples. Our pipeline requires a minimum of four samples, and this condition was satisfied in seventeen cases. In thirteen cases, all tumor samples were from the same kidney (unilateral). We obtained a flat phylogenetic
| Case number | Gender | Age at nephrectomy (months) | Local pathological stage | Histological subtype | Number of analyzed tumor samples | Germline DNA samples analyzed (NK: non-tumorous kidney; PBL: peripheral blood lymphocytes) | ADC analysis | Additional clinical features |
|-------------|--------|----------------------------|--------------------------|----------------------|----------------------------------|---------------------------------------------------------------------------------|-------------|-----------------------------|
| 1           | F      | 10                         | 1                        | Stromal              | 2                                | NK, PBL                                                                         | No          | Contralateral nephrogenic rest |
| 2           | M      | 18                         | 2                        | Mixed                | 3                                | None                                                                            | No          | Contralateral nephrogenic rest. Germline truncating WT1 mutation in exon 7 |
| 3           | M      | 32                         | 3                        | Mixed                | 3                                | PBL                                                                              | No          | Large tumor complicated by hypertensive cardiomyopathy. Surgery delayed due to anesthetic risk and chemotherapy extended until surgery at 10 weeks due to tumor enlargement. |
| 4           | F      | 58                         | 2                        | Blastemal            | 3                                | NK                                                                               | No          | Hemi hypertrophy             |
| 5           | M      | 48                         | 2                        | Regressive           | 2                                | NK                                                                               | No          | No                           |
| 6           | M      | 19                         | 3                        | Mixed                | 3                                | NK                                                                               | No          | Tetralogy of fallot          |
| 7           | F      | 10                         | 2                        | Mixed                | 3                                | NK                                                                               | Yes         | No                           |
| 8           | M      | 34                         | 2                        | Mixed                | 3                                | NK                                                                               | No          | Developmental delay. Constitutional gain of 22q11. 10 months before this partial nephrectomy: separate nephrogenic rest in same (left) kidney, confirmed on biopsy, and contralateral (right) regressive WT (after 9 weeks preoperative chemotherapy). Parents are first cousins |
| 9           | F      | 32                         | 1                        | Mixed                | 3                                | NK                                                                               | No          | Bilateral nephron-sparing surgery with tumor at margins bilaterally               |
| 10          | M      | 36                         | Right: 1                 | Right: mixed         | Left: 2                          | NK (left)                                                                       | No          | No                           |
| 11          | F      | 20                         | 1                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | Contralateral nephrogenic rest                                                |
| 12          | F      | 6                          | 1                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | Incidental diagnosis of WT during investigation of bilateral cryptorchidism. Germline WT1 substitution in exon 9 (His405Arg) |
| 13          | F      | 27                         | 3                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | No                           |
| 14          | M      | 12                         | 1                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | Bilateral nephron-sparing surgery with tumor at margins bilaterally               |
| 15          | M      | 14                         | 2                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | No                           |
| 16          | M      | 28                         | 3                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | No                           |
| 17          | F      | 5                          | 2                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | No                           |
| 18          | F      | 50                         | 3                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | Tumor rupture. Pulmonary metastases.                                           |
| 19          | F      | 42                         | 1                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | No                           |
| 20          | M      | 29                         | 1                        | Right: mixed         | Left: mixed                      | NK                                                                               | No          | Germline truncating WT1 mutation in exon 9 (associated with Denys-Drash syndrome) |
tree, where all samples contain the same clone, in Cases 3, 6 and 12. Cases 7 and 9 gave a linear tree, in which clones are derived from ‘ancestral’ clones sharing all prior evolutionary events. Branched evolution, in which more than one clone contains unique events compared with a common ancestor, was observed in the remaining seven unilateral tumors (1, 4, 8, 11, 13, 15, 18, 19).

Branched evolution is particularly well demonstrated by Cases 15 and 19, with marked branching that began early in the clonal evolution of the tumor. In Case 19 (Fig. 2), the most recent common ancestor clone observed in sample R4 shows 1q+ and 16q−. All other tumor samples display four additional events (+13, 16p+, +20, +X) and cluster separately from R5, R1 and R6. R2 contains a clone with two additional events (11q− and 13q13.1−) to R3. Interestingly, 13q13.1 loss removes a specific region encompassing BRCA2, after previous gain of the entire chromosome 13. R5, R1 and R6 share a +12 event that explains the distinct branching of the samples. However, further branching is seen in these samples as R5 contains two unique events (+6, +18), and R1 and R6 both exclusively share four events (+8, 3p25.3−, 14q22.2−23.3−, 22q13.31−). The clone present in R1 and R6 acquired three focal losses late in its evolution that encompass known cancer genes such as VHL, a tumor suppressor gene commonly mutated in clear cell renal cell carcinoma (ccRCC) (Gnarra et al., 1994), and SIX1, a gene that was recently reported to be recurrently mutated in WT (Wegert et al., 2015; Walz et al., 2015) (3p25.3− and 14q22.2−23.3−, respectively). Case 15 is even more diversified, with unique events in five of six samples (Supplementary Fig. 1).

3.3. Genetic Markers and Evolutionary Timing

3.3.1. Chromosome 11p15 Uniparental Disomy is Consistently an Early Event in WT Tumorigenesis

We observed 11p15 CNNLOH in 9 of 20 cases (Fig. 1). In six of these cases, LOH also involved the entire 11p13 region, and in Case 18, it involved part of the 11p13 region, including the WT1 locus; in Cases 4 and 8, 11p LOH did not involve 11p13. In all but Case 14, 11p15 CNNLOH was detected in all tumor samples, indicating that it occurred in a common ancestor of all observed clones in these tumors, and is thus a consistent early event in WT tumorigenesis. The observation that 11p15 CNNLOH was a truncal event in the evolution of these WTs was confirmed by phylogenetic analysis. In Case 14, 11p15 CNNLOH was called in just one of the two samples taken from this tumor, where it was the only CNA observed (Fig. 1). Visual inspection of the chromosome 11 BAF plots for this case shows that despite not being identified in one of the samples by our computational analysis, the event does appear to be present in a low fraction of cells (Supplementary Fig. 2). Therefore, Case 14 does not contradict the evidence from other cases that 11p15 CNNLOH is a consistently early event in WT tumorigenesis.

Fig. 1. Scope of intra-tumor copy number heterogeneity in 20 Wilms tumor cases. Case numbers, indicated on the y-axis, are split into three groups i) homogeneous, ii) bilateral and iii) heterogeneous. Chromosomes 1 to 22 and X are ordered on the x-axis from left to right, separated by vertical dashed lines. Cases are separated by solid black horizontal lines and samples from each case by dashed horizontal marks at each chromosome boundary. In the bilateral group, samples from contralateral kidneys are split by horizontal markers either side of the plot. Copy number states are displayed as white (expected copy number state), red (gain state), blue (loss state) and grey (expected copy number state with loss of heterozygosity).
Since chromosome 11p15 contains a cluster of imprinted genes, we performed MS-MLPA on the 11p15.2 locus (Scott et al., 2008a) in all cases with 11p CNNLOH, and in all tested samples found uniparental disomy (UPD), as indicated by hypermethylation of the H19 DMR (consistent with overexpression of the IGF2 oncogene) and hypomethylation of the KvDMR, i.e. a paternal pattern. DNA from adjacent normal kidney was available for eight of the nine cases with 11p15 CNNLOH, and in all eight cases, there was neither 11p CNNLOH nor abnormal methylation, indicating that the CNNLOH and methylation abnormalities were somatic events in the tumor cells.

In contrast, MS-MLPA on the cases without somatic 11p15 CNNLOH showed five cases (Cases 6, 10, 13, 17, 19) with hypermethylation of the H19 DMR only, with normal methylation of KvDMR. In these cases, the abnormal methylation pattern was also present in adjacent histologically normal kidney. Case 6 had known hemihypertrophy, but the other cases had not been diagnosed with Beckwith-Wiedemann syndrome, hemihypertrophy, abnormal growth, macroglossia, hypoglycemia or other tumors.

3.4. Gain of Chromosome 1q Shows Variable Timing

We observed 1q+ in eight patients (40%, Fig. 3; Supplementary Fig. 3). For seven of these we had multisampled the tumors in which we detected the change. 1q+ is present in all the tumor samples in three of these cases (Cases 11, 15, 19). In Cases 11 and 19 the same 1q+ is in all samples. In Case 15 there is 1q+ in all tumor samples (n = 6); however, there are several unique CNAs that affect this chromosome arm, thus displaying ITGH in the extent of 1q+ itself. In the remaining four cases, 1q+ is present in one of three (Cases 4 and 16), one of two (Case 16, right kidney tumor) and two of three (Case 20, right kidney tumor) samples. Thus, if we had single sampled our tumors, we would have only definitely identified three of seven tumors with 1q+; the probability of finding 1q+ in all seven tumors with single sampling is only 0.037. Indeed, the average probability of obtaining a negative single sample per tumor in our set of seven multi-sampled tumors with 1q+ is 0.31 (95% confidence interval is 0.08–0.54), despite 1q+ being present in all samples obtained from three of the seven positive tumors. The cumulative binomial distribution with p = 0.69 indicates that at least three samples per tumor must be studied in order to ensure that greater than 95% of tumors with 1q+ are detected.

In the four tumors which display 1q+ ITGH, 1q+ shows no preference in evolutionary timing. In Cases 9 and 16 chromosome 1q+ timing is indistinguishable from other copy number changes present in the tumor because samples from these tumors represent either the only clone with tumor-specific CNAs or have a copy number profile that is indistinguishable from matched germline DNA. In Cases 4 and 20 1q+
follows other CNAs (e.g. UPD of chromosome 11p in both cases) indicating that 1q+ evolved late. In contrast, in Cases 11 and 19 1q+ is a truncal event that may have occurred early in tumor evolution.

3.5. Rarer Biomarkers

Other WT biomarkers are difficult to assess without larger patient cohorts. Nevertheless, we observed 16q− in four tumors, three of which are multi-sampled (Fig. 1). Of these, Cases 18 and 19 show homogeneous 16q−, accompanied in Case 18 by homogeneous 1p− and 4q−. In Case 13, 16q− is apparently heterogeneous, but the change is present in the one sample that is from a tumor that probably originated independently in the same kidney (see below); therefore, we have no evidence to support 16q− as a heterogeneous event. Two cases contain a LOH event that affects the TP53 locus: homogeneous 17p CNNLOH in Case 18 and apparently heterogeneous loss of TP53 in Case 13 occurring in the single sample from the tumor with probable independent origin. Case 18 showed diffuse anaplasia on histological

Fig. 3. Chromosome 1q gain (1q+) is variably heterogeneous in eight cases. (A) Each sample is shown as a circle, filled in green if there is 1q+ and in white if there is not. Bilateral cases are split into contralateral samples with a horizontal black line. (B) Chromosome 1 copy number profiles for samples from Case 20. Log R ratios (y-axis) are shown for all chromosome 1 SNP probes, ordered by genomic position on the x-axis. Data points are coloured green if they correspond to copy number gains, and grey if not. Samples R1 and R4 display whole chromosomal arm gains.

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3.6. Spatially Separated Tumors

3.6.1. Bilateral Tumors

Five cases in our dataset are bilateral WTs, and in four of these both sides were sampled (Cases 10, 16, 17 and 20). Overall, bilateral tumors appear genetically distinct and probably arose independently. Cases 10 and 17 provide the clearest examples of the striking differences in copy number profiles between contralateral tumors. In Case 10 the only tumor-specific CNAs in the two samples taken from the left tumor are two focal deletions in chromosome 9, whereas the sample from the right tumor shows 1q+, +9, +12, +20, +X, 7p− and 16q−. In Case 17, the left tumor of Case 17 shows +7, +8, +12, +13, +16, +17, +22 and +X, while the right tumor shows no tumor-specific CNAs. In both these cases, the two contralateral tumors share no tumor-specific CNAs and are not heterogeneous within the individual kidneys. Case 16 only showed CNAs in one sample of two taken from the left tumor, thus making the right tumor heterogeneous. The left tumor sample of Case 16 did not have any CNAs.

Case 20 was heterogeneous within the right tumor, with 11p CNNLOH in all three samples but 1q+ in only two samples. The left tumor also contained 11p CNNLOH, but with a different centromeric boundary, indicating this may have been a separate unrelated event in the left tumor (Supplementary Fig. 2). However, we cannot formally exclude the possibility that the tumors in the two kidneys developed from a single clone with the shorter 11p CNNLOH, showing no further evolution in the left tumor and extension of 11p CNNLOH in the right kidney.

3.7. Evidence for Two Separate Tumors in the Same Mass

Case 13 presented as one tumor mass consisting of a larger middle nodule, consisting of mixed blastemal, epithelial and stromal elements, contiguous with a superior nodule composed of relatively well-differentiated epithelial structures, thus showing two contrasting WT phenotypes in the same apparent tumor. Four samples were obtained from the middle nodule and one from the superior nodule. The samples from the middle nodule were heterogeneous due to the presence or absence of +2, with homogeneous +6, +8, +9, +10, +12. The sample from the superior nodule contained 16q− and 17p−. Thus, the two nodules showed independent branching from germline DNA, suggesting that they arose independently, despite appearing as one contiguous tumor (Fig. 4).

3.7.1. Relation to Treatment Response

Case 8 also presented with a large intrarenal mass contiguous with a smaller nodule (Fig. 5A). Following neoadjuvant chemotherapy, however, the smaller nodule showed a greater shrinkage (78% vs. 42%, Fig. 5B) and increased ADC (0.71 to 1.71 vs. 0.73 to 1.32, Fig. 5C) than the main tumor mass, indicating a better response to chemotherapy (McDonald et al., 2010; Hales et al., 2015). Histologically, all areas of the tumor showed a mixed histology.

We obtained one sample from the smaller nodule (R1) and two from the larger mass (R3, R4); they show ITGH and phylogenetically, R4 represents a clonal ancestor of the other two samples. R3, also from the larger mass, evolved an additional Xp11.23−, while the smaller nodule developed 2q CNNLOH and a MYCN gain.

4. Discussion

Short-term outcomes in pediatric cancers are better than in adult cancers, but relapses are often unpredictable, and most current treatment protocols depend on cytotoxic agents with significant long-term complications (Reulen et al., 2010). There is a need to select better those patients who need more/less intensive therapy, and to increase the use of potentially safer targeted therapies. In WT, 1q+ is a promising genetic biomarker for such stratification, because it is both common and associated with a poorer outcome. Nevertheless, to date, assessment of its prognostic significance and suggested use in clinical trials has assumed that single tumor sampling is representative and sufficient. There is emerging evidence from a small number of recent studies that in WT and other pediatric solid tumors, a subset of cases show ITGH. We therefore hypothesized that ITGH in WT may involve prognostically significant CNA such as 1q+, and therefore we prospectively multi-sampled a series of twenty WT cases in our center. In parallel, we developed a custom analysis pipeline to detect reliably CNA and LOH events, compare them across multiple samples for each tumor, and infer evolutionary trajectories in order to provide a basis for understanding how ITGH arises. We found a remarkable range of evolutionary

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**Fig. 4.** Phylogenetic analysis reveals two genetically distinct Wilms tumors within a single mass in Case 13. The phylogenetic tree shows the evolutionary relationship between the normal kidney sample (NK, green node) and tumor samples (R1–5, red). The edges were drawn using the same rules as in Fig. 2. The tree shows that sample R5 is genetically distinct from R1–4, since their only common ancestral state corresponds to the normal kidney sample. The histology of the middle nodule, represented by R3, is triphasic, with blastemal, epithelial and stromal elements, whereas the superior nodule, represented by R5, is markedly different and composed exclusively of more mature epithelial elements (original magnification, ×100).
scenarios and variable ITGH in WT, including 1q+. Indeed, our data indicated that single sampling misses a significant proportion of cases with 1q+, and we estimated that to detect more than 95% of cases with 1q+, one would need to obtain at least three tumor samples per case.

We also found that 1q+ does not show preference in evolutionary timing—it may occur as an early or late event—which suggests that its oncogenic effect is independent of other genetic changes. Therefore, we suspect that 1q+ may have a similar effect on WT outcome regardless of whether it is homogeneous or heterogeneous in the primary tumor, and that current studies based on single tumor sampling may have underestimated its prognostic significance. Our findings clearly imply that future clinical trials in WT must take this heterogeneity into account and multi-sample each tumor or attempt to detect this change in circulating tumor DNA at a level that can interrogate subclonality.

In contrast, we find that somatic 11p15 CNNLOH, another common change in WT, is consistently an early event in WT tumorigenesis. Our finding therefore builds on previous observations of somatic 11p CNNLOH in WT precursor lesions (nephrogenic rests) (Charles et al., 1998). 11p15 CNNLOH is associated with several other pediatric small round cell tumors, and recently it was found as a recurrent lesion in the vast majority of pediatric adrenocortical tumors, also occurring as an early event preceding most point mutations (Pinto et al., 2015). These findings suggest that 11p15 CNNLOH may represent a common mechanism of tumorigenesis in a significant proportion of pediatric solid tumors, and its occurrence as an early event makes it a promising candidate for early detection of pediatric cancer by non-invasive screening for ctDNA in blood.

In those tumors without 11p15 CNNLOH we identified a subset of five cases with isolated hypermethylation of the H19 DMR, and this abnormality was also present in adjacent histologically normal kidney. In one of these five cases, there was hemihypertrophy, whereas in the other four cases there were no features to suggest Beckwith-Wiedemann syndrome. This finding is in keeping with previous reports of mosaic hypermethylation of the H19 DMR in a significant proportion of normal cells in cases of WT with this abnormality, even in the absence of other features of Beckwith-Wiedemann syndrome (Moultoun et al., 1994; Okamoto et al., 1997; Scott et al., 2008b). Indeed, the proportion and distribution of non-tumor cells with this WT-predisposing epimutation may at least in part underlie the expression and variability of the features of Beckwith-Wiedemann syndrome.
We have uncovered evidence of independent origins of two synchronous WT not only in bilateral cases but also within the same kidney containing an intra-tumoral nodule with divergent histology. While the presence of multicentric tumors in the same kidney is a recognized feature in 5%–10% of WT, in Case 13 the nodule with divergent histology was contiguous with the main tumor mass. Multicentric WT may thus be under-recognized and therefore not treated appropriately. Furthermore, under current diagnostic criteria, the relative proportions of blastemal, epithelial and stromal elements are used to stratify WT into low/intermediate/high-risk categories, with an underlying assumption that such structures are all derived from the same tumor. However, this practice needs to be refined to take into account multiple tumors, of independent origins, within the same overall mass as well as multicentric and bilateral WT.

Our findings on rarer biomarkers are more difficult to interpret in the absence of a larger multi-sampled tumor cohort. Nevertheless, our findings on 16q — highlight the importance of interpreting ITGH in the context of tumor evolution: in our series, 16q — is apparently heterogeneous only in Case 13, but it is erroneous to interpret this as evidence of 16q — ITGH, since it is present in the one sample from the smaller nodule that we showed arose independently of the remaining tumor mass. In the case of another biomarker, MYCN gain, we were able to relate subclonal acquisition of this change to a significantly better response to chemotherapy (as assessed on diffusion-weighted MR imaging before and after preoperative chemotherapy). MYCN gain may be expected to be associated with more rapid cell proliferation and therefore greater sensitivity to cytotoxic chemotherapy, and this may explain our finding. More generally, we have shown that it is feasible to integrate phylogenetic tumor analysis with monitoring of treatment response by imaging, provided that the imaging analysis is used as a guide to tumor sampling, in addition to current standard histological sampling.

Taken together, our findings in WT show unpredictable and clinically significant genetic heterogeneity that requires tumor multisampling for its detection, and assessment of tumor evolutionary trajectories for its interpretation. The custom analysis pipeline that we developed for this project may be easily applied to similar data from other multi-sampled tumors, and we are also extending it to integrate single nucleotide variants and small indels, which are typically detected in sequencing studies. Our findings have major implications for planning biomarker sampling strategies in future clinical trials for WT, and possibly other pediatric solid tumors.

Author Contributions

GDC, JRA, BM, NJS, KPJ, RDW and WM designed the study. JRA obtained and organized clinical, radiological imaging and macropathological data. TC extracted DNA and carried out molecular analyses. ÖEO performed radiological imaging analyses. SJP and WM reviewed histology and WM took photomicrographs. GDC, BM, NJS, KPJ, NML, RDW and WM supervised the project. GDC prepared the figures. GDC, JRA, BM, NJS, KPJ, NML, RDW and WM wrote the paper.

Declaration of Interests

None.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.ebior.2016.05.029.

References

Charles, A.K., Brown, K.W., Berry, P.J., 1998. Microdissecting the genetic events in nephrogenic rests and Wilms’ tumor development. Am. J. Pathol. 153, 991–1000.

de Bruin, E.C., McGranahan, N., Mitter, R., et al., 2014. Spatial and temporal diversity in geno
sic instability processes defines lung cancer evolution. Science 346, 251–256.

Diskin, S.J., Li, M., Hou, C., et al., 2008. Adjustment of genomic waves in signal intensities from whole-genome SNP genotyping platforms. Nucleic Acids Res. 36, e126.

Gentleman, R.C., Carey, V.J., Bates, D.M., et al., 2004. Bioconductor: open software development for computational biology and bioinformatics. Genome Biol. 5, R80.

Gelger, M., Rowan, A.J., Horswell, S., et al., 2012. Intratumoral heterogeneity and branched evolution revealed by multiregion sequencing. N. Engl. J. Med. 366, 883–892.

Gelger, M., Horswell, S., Larkin, J., et al., 2014. Genomic architecture and evolution of clear cell renal cell carcinomas defined by multiregion sequencing. Nat. Genet. 46, 225–233.

Gnarra, J.R., Toy, K., Weng, Y., et al., 1994. Mutations of the VHL tumour suppressor gene in renal carcinoma. Nat. Genet. 7, 85–90.

Gratias, E.J., Jennings, L.J., Anderson, J.R., Dome, J.S., Grundy, P., Perlman, E.J., 2013. Gain of 1q is associated with inferior event-free and overall survival in patients with favorable histology Wilms tumor: a report from the Children’s Oncology Group. Cancer 119, 3887–3894.

Graves, M., Mäley, C.C., 2012. Clonal evolution in cancer. Nature 481, 306–311.

Grundy, P.E., Breslow, N.E., Li, S., et al., 2005. Loss of heterozygosity for chromosomes 1p and 16q is an adverse prognostic factor in favorable-histology Wilms tumor: a report from the National Wilms Tumor Study Group. J. Clin. Oncol. 23, 7312–7321.

Hales, P.W., Olsen, B.E., Sebire, N.J., Pritchard-Jones, K., Clark, C.A., 2015. A multi-Caucasian model for apparent diffusion coefficient histogram analysis of Wilms tumor subtype and response to chemotherapy. NMR Biomed. 28, 948–957.

McDonald, K., Sebire, N.J., Anderson, J., Olsen, B.E., 2010. Patterns of shift in ADC distributions in abdominal tumours during chemotherapy—feasibility study. Pediatr. Radiol. 41, 99–106.

Mengelberger, L.H., Karlsson, J., Lindgren, D., et al., 2015. Intratumoral genome diversity parallels progression and predicts outcome in pediatric cancer. Nat. Commun. 6, 6125.

Moulton, T., Crenshaw, T., Hao, Y., et al., 1994. Epigenetic lesions at the H19 locus in Wilms’ tumour patients. Nat. Genet. 7, 440–447.

Natrajan, R., Williams, R.D., Hing, S.N., et al., 2006. Array CGH profiling of favourable histology Wilms tumours reveals novel gains and losses associated with relapse. J. Pathol. 210, 49–58.

Newell, P.C., 1976. The clonal evolution of tumor cell populations. Science 194, 23–28.

Okamoto, K., Morison, L.M., Taniguchi, T., Reeve, A.E., 1997. Epigenetic changes at the in
sulin-like growth factor II/H19 locus in developing kidney is an early event in Wilms tumorigenesis. Proc. Natl. Acad. Sci. U. S. A. 94, 5367–5371.

Pinto, E.M., Chen, X., Easton, J., et al., 2015. Genomic landscape of paediatric adrenocortical tumours. Nat. Commun. 6, 6302.

Popov, I., spanjic, G.M., Sebire, N.J., et al., 2013. Bilateral Wilms tumours with TP53-relat
ded anaplasia. Pediatr. Dev. Pathol. 16, 217–223.

Pritchard-Jones, K., Bergeron, C., de Camargo, B., et al., 2015. Omission of doxorubicin from the treatment of stage II-III, intermediate-risk Wilms’ tumour (SIOP WT 2001): an open-label, non-inferiority, randomised controlled trial. Lancet 386, 1156–1164.

Remke, M., Cavalli, F., Morriess, A., et al., 2015. Integrated genomics elucidates relative spatial homogeneity of embryonal brain tumors. Mol. Cell Pediatr. 2, A10.

Reulen, R.C., Winter, D.L., Frohisher, C., et al., 2010. Long-term cause-specific mortality among survivors of childhood cancer. JAMA 304, 172–179.

Schlieermann, G., Michon, J., Ribeiro, A., et al., 2011. Segmental chromosomal alterations lead to higher risk of relapse in infants with MYCN-non-amplified localised unresectable/disseminated neuroblastoma (a SIOPEN collaborative study). Br. J. Cancer 105, 1940–1948.

Schwarz, R.F., Triath, A., Sipos, B., Brenton, J.D., Goldman, N., Markowitz, F., 2014. Phyloge
etic quantification of intra-tumour heterogeneity. PLoS Comput. Biol. 10, e1003535.

Scott, R.H., Douglas, J., Baskcomb, L., et al., 2008a. Methylation-specific multiplex ligation dependent probe amplification (MS-MLPA) robustly detects and distinguishes 1p15 abnormalities associated with overgrowth and growth retardation. J. Med. Genet. 45, 106–113.
Scott, R.H., Douglas, J., Baskcomb, L., et al., 2008b. Constitutional 11p15 abnormalities, including heritable imprinting center mutations, cause nonsyndromic Wilms tumor. Nat. Genet. 40, 1329–1334.
Segers, H., van den Heuvel-Eibrink, M.M., Williams, R.D., et al., 2013. Gain of 1q is a marker of poor prognosis in Wilms’ tumors. Genes Chromosom. Cancer 52, 1065–1074.
van de Wiel, M.A., van Wieringen, W.N., 2007. CGHregions: dimension reduction for array CGH data with minimal information loss. Cancer Informat.http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2675846/
van de Wiel, M.A., Kim, K.J., Vosse, S.J., van Wieringen, W.N., Wilting, S.M., Ylstra, B., 2007. CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics 23, 892–894.
Vogelstein, B., Papadopoulos, N., Velculescu, V.E., Zhou, S., Diaz Jr., L.A., Kinzler, K.W., 2013. Cancer genome landscapes. Science 339, 1546–1558.
Vujanić, G.M., Sandstedt, B., Harms, D., Kelsey, A., Leuschner, I., de Kraker, J., 2002. Revised International Society of Paediatric Oncology (SIOP) working classification of renal tumors of childhood. Med. Pediatr. Oncol. 38, 79–82.
Walz, A.L., Ooms, A., Gadd, S., et al., 2015. Recurrent DGCR8, DROSHA, and SIX homeodomain mutations in favorable histology Wilms tumors. Cancer Cell 27, 286–297.
Wegert, J., Wittmann, S., Leuschner, I., Geissinger, E., Graf, N., Geisler, M., 2009. WTX inactivation is a frequent, but late event in Wilms tumors without apparent clinical impact. Genes Chromosom. Cancer 48, 1102–1111.
Wegert, J., Ishaque, N., Vardapour, R., et al., 2015. Mutations in the SIX1/2 pathway and the DROSHA/DGCR8 mRNA microprocessor complex underlie high-risk blastemal type Wilms tumors. Cancer Cell 27, 298–311.
Williams, R.D., Chagtai, T., Alcaide-German, M., et al., 2015. Multiple mechanisms of MYCN dysregulation in Wilms tumour. Oncotarget 6, 7232–7243.