Pediatric Extra Renal Renal Cellular Carcinoma with Xp11 Translocation Responding to Sunitinib, Everolimus and Metronomic Chemotherapy

István Szegedi*, Gábor Méhes, Imre Gáspár, Tamás Csonka, István Csizy and Csongor Kiss
Department of Pediatrics, University of Debrecen, Hungary

Abstract

Background: Fewer than 2% of renal cell carcinomas (RCC) develop in childhood, making up less than 0.1-0.3% of all pediatric tumors and 2.6% of renal neoplasms. The 2004 WHO classification defined RCC with Xp11.2 aberration as a separate entity accumulating characteristically in childhood. Ectopic development of pediatric renal neoplasms is rare and there are no data in the literature about the extra renal origin of RCC.

Case Presentation: We describe the case of a 16-year-old boy with an extra renal RCC showing Xp11.2 translocation demonstrated by FISH. The tumor exhibited a favourable response to a combination of targeted treatment with sunitinib and everolimus in association with metronomic chemotherapy.

Conclusion: The novelty of the reported case, in addition to the extra renal origin of the tumor, is the favourable response to a combination of sunitinib, everolimus and metronomic chemotherapy resulted in a major partial control of advanced RCC accompanied by a 15 months survival from diagnosis without major toxicities.

Keywords: Renal cell carcinoma; pediatric; Extra renal; Xp11 translocation; Targeted therapy; Metronomic treatment

Background

Renal cell carcinoma (RCC) is a rare form of childhood cancer representing fewer than 0.1-0.3% of all tumors and 2.6% of renal neoplasms in the pediatric population. Less than 2% of all RCC cases occurs in pediatric patients, with a highest incidence between 15-19 years [1-2]. The 2004 WHO classification defined RCC with Xp11.2 aberration as a separate entity accumulating characteristically in childhood. RCC associated with Xp11.2 translocation characteristically affects children; however, adult cases may outnumber pediatric cases due to a much higher overall incidence in the adult population [3-4]. The most frequent pediatric renal neoplasm is Wilms tumor which rarely may originate from an extra renal ectopic nephrogenic rest [5-6]. There is no data, however, in the literature about the extra renal origin of RCC with classical clear cell morphology. Targeted therapies with broad spectrum tyrosine kinas inhibitors (TKI), agents interfering with the effects of vascular endothelial growth factor (VEGF) and inhibitors of the Mammalian Target of Rapamycin (MTOR) have been shown superior to either immunotherapy or to placebo treatment in randomized phase III trials designed for adult patients with RCC [7]. The above agents have not been registered for use in children and pediatric experiences with the treatment of RCC are restricted to sporadic case reports only [8]. Combined oral metronomic bio differentiating anti angio genetic treatment /COMBAT/ proved promising in a series of advanced refractory pediatric malignancies [9]. However, there are only limited data on the efficacy and safety of metronomic therapy in RCC [10]. Here we report, for the first time, on the application of the multi targeted oral TKI sunitinib in combination with the MTOR inhibitor everolimus and metronomic chemotherapy resulting in major partial response in a 16-year-old boy with an extensive extra renal RCC.

Case Presentation

A 16-year-old Caucasian male was referred to the Department of Pediatric Hematology-Oncology, University of Debrecen, in June, 2010 because of lower limb pain for the last six months, fatigue, anorexia, and significant weight loss. On referral he was in a poor general condition, was significantly underweight (height: 175 cm; 50-75 percentile; weight: 45 kg; < 3 percentile). Abdominal
ultra sonography showed a 5.5 cm large in homogeneous lesion at the left suprarenal region containing areas of calcification. Contrast enhanced abdominal computed tomography scan confirmed the same mass, which was completely separated from the left kidney (Figure 1A). There were enlarged paraaortic lymph nodes present, and an additional destructive 10 cm large lesion on the right side of the pubic bone, with wide muscular infiltration, and an epidural mass with intra spinal invasion at the L2 region. Magnetic resonance imaging showed a left par vertebral necrotic mass spreading behind the left renal vein and artery, a 10 mm enhancing mass at the corpus of ThVIII, a compression of ThXII resulting in the narrowing of the spinal canal, and another mass compressing the cone at the level of LII (Figure 1B). In addition, there was an infiltration of the sacral bones in both sides. Tc-99m metilen-diphosphonat bone scan revealed an additional right sided infiltration of the hip joint. F-131 MIBG scan did not show enhancement of the tracer. Initial laboratory tests revealed moderate anaemia (86g/L), a normal Neuron Specific Enolase (NSE) level and elevated plasma noradrenalin and urinary dopamine levels. Neurosurgery was performed to achieve decompression at the level of ThXII. The suprarenal location of the primary lesion and the elevated catecholamine level prompted us to start an urgent chemotherapeutic treatment according to the SIOP High Risk Neuroblastoma protocol [11]. Despite treatment, progressive disease was seen. In the meantime, histological evaluation revealed a necrotic and infiltrative clear cell carcinoma, which was positive for Oil-red-O, CD10, Bcl2, NSE and negative for Pan-Cytokeratin (PCK), GFAP, chromogranin, synaptophysin, desmin, inhibit and nest in, a typical morphological and immune phenotypic pattern suggesting the diagnosis of RCC with Xp11 translocation (Figure 2). Unresponsiveness to conventional chemotherapy was consistent with the diagnosis of RCC therefore; we started sunitinib treatment (1x50mg in 4 week-cycles, 6 weekly) and oral cyclophosphamide (2x25 mg/day) and isotretinoin as a part of COMBAT metronomic therapy [9]. The patient experienced an 8-months-long period of stable disease. Due to progression in June 2011 mTOR inhibitor everolimus has been added to the ongoing therapy (2x10 mg/day). After a week of treatment there was a prominent decrease in tumor volume as checked by physical examination with an almost complete disappearance of the huge inguinal resistance. In August, 2011 a new histological sample was taken from this inguinal region. Microscopy showed a necrotic and haemorrhagic tissue mass without signs of viable RCC infiltration. In September, 2011 a Tc 99m bone-scan revealed major regression of the former metastatic regions without signs of osteoblastic activation. Later in September, 2011 the patient developed an excessive pneumonia and succumbed to death despite combined antibiotic treatment and mechanical ventilation. Autopsy findings were in agreement with the previous images including the suprarenal localisation of the primary lesion which was independent from the kidney. (Figure 1 C, D). Post-mortem histology confirmed the diagnosis of clear cell cancer equivalent in every regard of RCC. Residual tumor masses contained large areas of necrosis and hemorrhage intermixed with focal scarring. Intact tumor was present only in small foci.

Methods

Histological examination was performed following fixation in 4.0% buffered formaldehyde and paraffin embedding as usual. Four-micrometer-thick sections were prepared and stained with hematoxilin-eosin. Immunohistochemistry was carried out with the Bond Max immune stainer (Leica, Wetzlar, Germany) using a panel of antibodies for PCK, CD10, Bcl2, NSE, GFAP, chromogranin, synaptophysin, desmin, inhibit and nest in, a typical morphological and immune phenotypic pattern suggesting the diagnosis of RCC with Xp11 translocation (Figure 2). Unresponsiveness to conventional chemotherapy was consistent with the diagnosis of RCC therefore; we started sunitinib treatment (1x50mg in 4 week-cycles, 6 weekly) and a complex bone supportive therapy (sodium clodronate, vitamin D) in September, 2010. Later, sunitinib treatment was supplemented with oral cyclophosphamide (2x25 mg/day) and isotretinoin as a part of COMBAT metronomic therapy [9]. The patient experienced an 8-months-long period of stable disease. Due to progression in June 2011 mTOR inhibitor everolimus has been added to the ongoing therapy (2x10 mg/day). After a week of treatment there was a prominent decrease in tumor volume as checked by physical examination with an almost complete disappearance of the huge inguinal resistance. In August, 2011 a new histological sample was taken from this inguinal region. Microscopy showed a necrotic and haemorrhagic tissue mass without signs of viable RCC infiltration. In September, 2011 a Tc 99m bone-scan revealed major regression of the former metastatic regions without signs of osteoblastic activation. Later in September, 2011 the patient developed an excessive pneumonia and succumbed to death despite combined antibiotic treatment and mechanical ventilation. Autopsy findings were in agreement with the previous images including the suprarenal localisation of the primary lesion which was independent from the kidney. (Figure 1 C, D). Post-mortem histology confirmed the diagnosis of clear cell cancer equivalent in every regard of RCC. Residual tumor masses contained large areas of necrosis and hemorrhage intermixed with focal scarring. Intact tumor was present only in small foci.

Methods

Histological examination was performed following fixation in 4.0% buffered formaldehyde and paraffin embedding as usual. Four-micrometer-thick sections were prepared and stained with hematoxilin-eosin. Immunohistochemistry was carried out with the Bond Max immune stainer (Leica, Wetzlar, Germany) using a panel of antibodies for PCK, CD10, Bcl2, NSE, GFAP, chromogranin, synaptophysin, desmin, inhibit and nest in, a typical morphological and immune phenotypic pattern suggesting the diagnosis of RCC with Xp11 translocation (Figure 2). Unresponsiveness to conventional chemotherapy was consistent with the diagnosis of RCC therefore; we started sunitinib treatment (1x50mg in 4 week-cycles, 6 weekly) and a complex bone supportive therapy (sodium clodronate, vitamin D) in September, 2010. Later, sunitinib treatment was supplemented with oral cyclophosphamide (2x25 mg/day) and isotretinoin as a part of COMBAT metronomic therapy [9]. The patient experienced an 8-months-long period of stable disease. Due to progression in June 2011 mTOR inhibitor everolimus has been added to the ongoing therapy (2x10 mg/day). After a week of treatment there was a prominent decrease in tumor volume as checked by physical examination with an almost complete disappearance of the huge inguinal resistance. In August, 2011 a new histological sample was taken from this inguinal region. Microscopy showed a necrotic and haemorrhagic tissue mass without signs of viable RCC infiltration. In September, 2011 a Tc 99m bone-scan revealed major regression of the former metastatic regions without signs of osteoblastic activation. Later in September, 2011 the patient developed an excessive pneumonia and succumbed to death despite combined antibiotic treatment and mechanical ventilation. Autopsy findings were in agreement with the previous images including the suprarenal localisation of the primary lesion which was independent from the kidney. (Figure 1 C, D). Post-mortem histology confirmed the diagnosis of clear cell cancer equivalent in every regard of RCC. Residual tumor masses contained large areas of necrosis and hemorrhage intermixed with focal scarring. Intact tumor was present only in small foci.
the TFE3 transcription factor gene region at Xp11. A break was defined as a separate red (centromeric) and green (telomeric) signals demonstrating TFE3 rearrangement in tumor cells in contrast to red-green signals representing the normal TEF3 gene. 72/100 cell nuclei were identified with the break apart FISH signal pattern.

Conclusions

In spite of major advances in therapy, metastatic RCC is still considered an incurable disease. In adults, 25-30% of patients present with metastases at time of diagnosis [12]. In contrast to many other forms of cancer, introduction of combined chemotherapy and immunotherapy did not change the grave prognosis of RCC which proved a constitutionally chemoresistant malignant disease. Targeted therapy aimed at involved signalling pathways has become first line treatment [12]. Treatments with multi kinase inhibitors and MTOR inhibitors resulted usually in Disease Stabilization (SD) and elicited partial responses, whereas complete responses were rarely reported [13]. Several ongoing clinical studies investigate the efficacy of different combinations of targeted agents; however, none of the experimental combinations have yet been introduced in routine clinical practice. Experiences with RCC occurring in children are still exceptional. COMBAT is a feasible and effective treatment option for patients with refractory/relapsing malignancies characterized by a low toxicity profile; however, it has not been applied in pediatric RCC [9]. In this report we presented an aggressive extra renal clear cell tumor characterized by Xp11 translocation in a 16-year old boy, which was equivalent to RCC in all aspects [4]. To the best of our knowledge this is the first extra renal case of RCC, regardless of the age, that has been reported in the literature. Detailed morphological analysis could not identify any other tissue component to the clear cell component which was moderately differentiated forming tubular arrangement (Figure 2A). The characteristic immune profile of the cells was rather homogenous inducing CD10 and bcl-2 immune positivity next to negativity for general mesenchymal and neurogenic markers (Figure 2B). FISH analysis performed on the few residual tumor cell nests demonstrated the duplication of the X chromosome due to two copies of both the green and the red FISH signals. The characteristic break apart Xp11 translocation pattern was stated on both (supernumerary) X chromosomes (Figure 3) as green and red signals completely separated in the majority (72%) of cell nuclei. The novelty of the reported case, in addition to the extra renal origin of the tumor, is the favourable response to a combination of sunitinib, everolimus and metronomic chemotherapeutic therapy. The oral multi targeted receptor tyrosine kinase inhibitor sunitinib displays a 31% objective response rate in metastatic RCC patients resulting in a median progression free survival from 5-11 months [14]. For patients with advanced, metastatic RCC who progressed on VEGF receptor tyrosine kinase inhibitor therapy, the MTOR inhibitor everolimus has been shown to prolong progression free survival vs. placebo from 1.9 months to 4.9 months. Unfortunately, combination of everolimus and sunitinib was associated with significant acute and chronic toxicities at standard doses and showed infrequent efficacy in clear cell RCC in an adult phase I trial [7]. Here we reported that combined use of targeted therapies combined with metronomic type of cytostatic treatment resulted in a major partial control of advanced RCC accompanied by a 15 months survival from diagnosis. No major toxicities were observed. The presented case may suggest possible merits of combining TKI-s and MTOR inhibitors with COMBAT in metastatic RCC. The death of the patient was independent from tumor progression as proven by the autopsy findings.

Consent

Written informed consent was obtained from the patient’s mother for publication of this Case report and any accompanying images. A copy of the written consent is available for review by the Editor of this journal.

List of Abbreviations

RCC – renal cell carcinoma; WHO – World Health Organization; FISH – fluorescence in situ hybridization; TKI – tyrosine kinase inhibitor; VEGF – vascular endothelial growth factor; MTOR – mammalian target of rapamycin; COMBAT – combined oral metronomic bio differentiating anti angio genetic treatment; MiBG – metiodibenzy-l-guanidine; NSE – neuron specific enolase; SIOP-International Society of Pediatric Oncology; CD10 – cell differentiation antigen; Bcl2 – B-cell lymphoma 2; PCK – pan-cytokeratin; GFAP – glial fibrillary acidic protein; TFE3 – transcription factor E3; SD – stable disease.
Contributors' Statement

István Szegedi: Dr. Szegedi designed and organized the main diagnostic procedure of the case, planned and conducted the particular treatment steps and conceptualized and designed this case report, drafted the initial manuscript, and approved the final manuscript as submitted.

Gábor Méhes: Prof. Méhes coordinated the histological and molecular diagnostic investigations and the autopsy, and reviewed and critically revised the manuscript, and approved the final manuscript as submitted.

Imre Gaspar: Dr. Gaspar helped to carry out the main steps of the treatment, and reviewed the manuscript, and approved the final manuscript as submitted.

Tamás Csonka: Dr. Csonka organized and carried out the histological, molecular diagnostic investigations and the autopsy, and reviewed the manuscript, and approved the final manuscript as submitted.

István Csízy: Dr. Csízy carried out the surgical invasive procedures and performed all the sampling of the patient, and reviewed the manuscript, and approved the final manuscript as submitted.

Csongor Kiss: Prof. Kiss supervised the diagnostic and the therapeutic steps, critically reviewed and revised the manuscript, and approved the final manuscript as submitted.

Acknowledgement

This work was supported by TÁMOP 4.2.2.A-11/1/KONV-2012-0025 project which are implemented through the New Hungary Development Plan, co-financed by the European Social Fund.

References