Description of longitudinal tumor evolution in a case of multiply relapsed clear cell sarcoma of the kidney

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Abstract
Background: Clear cell sarcoma of the kidney (CCSK) is the second most common pediatric renal tumor.
Case: A 2-year-old boy was diagnosed with CCSK, which relapsed four times until he yielded to the disease at the age of 7 years. To characterize the longitudinal genetic alterations occurring in the present case, we performed targeted-capture sequencing by pediatric solid tumors panel (381 genes) for longitudinally sampled tumors, including autopsy samples of metastasis. Internal tandem duplication of BCOR (BCOR-ITD) was the only truncal mutation, confirming the previously reported role of BCOR-ITD in CCSK.
Conclusion: Acquisition of additional mutations along tumor relapses and detection of metastasis-specific mutations were reminiscent of the tumor progression and therapeutic resistance of this case, leading to clonal selection and a dismal fate.

Keywords
BCOR-ITD, clear cell sarcoma of the kidney, clonal evolution, relapse

1 | INTRODUCTION

Clear cell sarcoma of the kidney (CCSK) makes up 4% of all primary renal malignancies in children.1 Although its prognosis has significantly improved recently,1–5 the relapse rate is still high and the prognosis of patients with relapses is extremely poor.6 Internal tandem duplication of BCOR (BCOR-ITD)7,8 YWHAE-NUTM2B/E fusion,9,10 and BCOR-CCNB3 fusion11 have been reported in CCSK cases; however, the genetic mechanisms associated with tumor recurrence and metastasis are still poorly understood. In this study, we assessed the genetic mechanisms of CCSK recurrence and metastasis by analyzing longitudinally sampled tumors extracted from a single case: specimens at diagnosis, each relapse, and autopsy.
A 2-year-old boy presented with a 2-month history of abdominal distention. At the initial diagnosis, a tumor \((17 \times 14 \times 11 \text{ cm})\) in the right kidney without metastasis (Stage-I) was detected by enhanced computed tomography (CT) (Figure S1). The patient underwent a right radical nephrectomy without lymph node dissection (Figure 1), and pathological findings led to a diagnosis of CCSK. He received chemotherapy consisting of actinomycin D, vincristine, and doxorubicin (DD-4A) with radiotherapy \((10.8\text{ Gy/6Fr})\) and this combination therapy successfully induced complete remission. However, 4 months after the completion of treatment, a first relapse occurred at his left lung. He received six courses of chemotherapy consisting of ifosfamide, carboplatin, and etoposide (ICE) with radiotherapy \((12\text{ Gy/8Fr})\) after a left upper lobectomy, and then achieved a second remission. At the age of 5 years, a second relapse developed at the para-aortic lymph nodes. A total excision was done after three courses of chemotherapy consisting of cyclophosphamide, vincristine, pirarubicin, and cisplatin. Histologically, viable tumor cells remained with fibrosis and mucoid substances (Supplementary Material S1). Postoperative chemoradiotherapy induced a third remission. At the age of 6 years, he suffered from a third relapse at the para-aortic lymph nodes, the same location as the second relapse. Chemotherapy consisting of ifosfamide, etoposide, vincristine, doxorubicin, and cyclophosphamide resulted in a poor therapeutic response. The patient underwent a subtotal excision of the lesion. Pathological examination of resected specimen detected viable tumor cells among therapy-related necrosis and fibrosis (Supplementary material S1). High dose chemotherapy (busulfan and melpharan) with autologous hematopoietic stem cell transplantation (aHSCT) induced a fourth remission; however, 4 months later, a fourth relapse occurred at the ascending colon, small intestine, duodenum, pancreas, and right hepatic lobe, and he yielded to the advanced disease at the age of 7 years.

### RESULTS

After obtaining informed consent from his parents, DNA was extracted from the peripheral blood and biopsy samples obtained at initial diagnosis, each relapse (second relapse; R2, third relapse; R3, fourth relapse; R4) and autopsy (abdominal lymph node; LN, liver; Liver) except for the sample at first relapse. Direct sequencing identified BCOR-ITD \((c.5526\text{_}5621\text{dup})\) at all the analyzed samples (Figure S2). SNP array analysis did not reveal any copy number alterations (CNAs) at the diagnosis whereas consistent CNAs on chromosomes 1q, 7q, and 15q were detected in R2 through R4 as well as LN and liver (Figure 2). Each sample possessed several sample specific CNAs. Moreover, we detected a focal loss at chromosome 9q32-33 in the samples of R2 through autopsy, where only the tumor suppressor gene BRIP1 was located (Figure 3A). We further analyzed other genetic alterations using targeted-capture sequencing (TCS) for 381 genes relating to pediatric solid tumors. Somatic mutations...
were filtered using the peripheral blood sample as germline control. Candidate somatic mutations located in exonic regions were further filtered by excluding variants: (a) with a VAF < 0.04; (b) listed in SNP databases; and (c) synonymous single-nucleotide variants, according to our previous report.\textsuperscript{12} Finally, 13 mutations were detected and classified into 7 clones based on the detected samples and VAF (Table 1 and Figure 3B). Although validation of the candidate mutations was not performed in this study, our sequence depth (mean 600-714) combined with EB call (Empirical Bayesian Mutation calling) analysis\textsuperscript{13} in Genomon software (https://github.com/Genomon-Project/GenomonPipeline) was estimated to be sufficient to rule out sequencing artifacts. Only the \textit{BCOR}-ITD was included in clone 1, whereas three clade mutations (\textit{FLT1}, \textit{JAK2}, and \textit{MYH7}) were additionally detected in relapsed tumor samples (clones 2, 3). The other mutations (\textit{PRKRIR}, \textit{KMT2D}, \textit{TSHR}, \textit{CDK6}, \textit{IL6ST}, \textit{NF1}, \textit{KDM6A}, \textit{OBSCN}, and \textit{SYNE1}) were clone specific (clones 4-7). The number of detected mutations increased during each tumor recurrence. All R3 mutations (clones 1-3) were detected upon metastatic samples from autopsy, suggesting the accumulation of metastasis site-specific mutations during tumor progression. Importantly, the newly acquired mutations at each relapse were not detected even as a minor clone in the previous samples, though variants with a VAF < 0.02 were not detectable due to our mutation calling method (Supplementary Material S2). We analyzed the clonal evolution of the present case based on the VAF of

\begin{figure}
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\caption{Results of SNP array analysis in each sample. Colorful dots at the top represent signal of each probe (raw data). Red and green lines at the bottom show allele-specific copy number, respectively. Consistent copy number alterations on chromosomes 1q, 7q, and 15q were detected. D, initial diagnosis; LN, lymph node; R2, second relapse; R3, third relapse; R4, fourth relapse.}
\end{figure}
each mutation and constructed a node-based phylogenetic tree using a bootstrap resampling technique implemented in ClonEvol.\textsuperscript{14,15} Longitudinal sequencing analysis at different time points allowed us to delineate the clonal history of tumor cells, suggesting that each relapse developed based on the preexisting tumors (Figure 3C).

4 | DISCUSSION

In this study, we presented a case of CCSK, which underwent four relapses. The first relapse occurred in the lung. This is consistent with the previous reports that the lung was the most common site of relapse in patients treated with DD-4A,\textsuperscript{4,5} although the present case relapsed earlier (4 vs 24 months).\textsuperscript{1,3} The optimal treatment of relapsed CCSK has not been established as yet. ICE chemotherapy, a salvage regimen for recurrent CCSK,\textsuperscript{6,16} resulted in the achievement of the second remission, however, the second relapse developed. Due to the refractory disease, we performed high dose chemotherapy with aHSCT rescue though its effectiveness is unclear.\textsuperscript{6,16} Pathological analysis of post-treatment specimens revealed a partial response, consistent with clinical therapeutic resistance. Because treatment options for relapsed CCSK are limited and survival is poor, we retrospectively...
analyzed this case in terms of genomics for the possibility of new therapeutic strategies.

In this study, the same BCOR-ITD alteration was observed from diagnosis right up to autopsy with very low tumor mutation burden, suggesting a strong impact not only on CCSK tumorigenesis as previously reported\(^7,8\) but also on its relapse. Meanwhile, we could not observe truncal mutations that can be targetable with available FDA-approved drugs.

Accumulation of the genetic abnormalities during tumor progression might be associated with the therapeutic resistance in this case. BRIP1, lost in the samples of R2 through autopsy, is a tumor suppressor gene, which regulates the G1/S checkpoint and cell cycle\(^17\) and its dysregulation is associated with tumor recurrence in other carcinomas.\(^18\) Our clonal analysis of the metastatic samples suggested that relapsed tumors were derived from minor clones of the preceding tumors, which had survived chemoradiotherapy and thrived (Figure 3C). Moreover, multisampled extractions at autopsy revealed site-specific mutations of metastasis in addition to truncal mutations of R3 sample, which suggests the presence of subclones with site-specific mutations in each metastases site of the previous samples. Among them, a subclone with selective advantage might relapse as the dominant clone. This is supported by the result that the newly acquired mutations were not detected even as a minor clone in the previous primary samples. However, our study is based on a single case and functions of the newly acquired genetic alterations remain unclear. In addition, our analysis in the present study was limited to genomic abnormalities, not including other various factors such as epigenetic abnormalities and tumor microenvironment. Further studies with more samples will help to detect common mechanisms of relapse and therapy resistance in CCSK, enabling appropriate therapy for relapsed cases considering genetic status.

Altogether, we failed to find targetable lesions, while our study indicated the importance of BCOR-ITD in both CCSK tumorigenesis and relapse. The genetic alterations acquired during disease progression might contribute to clonal selection and therapeutic resistance in the present case though further studies with more samples are warranted.

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### CONFLICT OF INTEREST

The authors declare no conflicts of interest.

### AUTHOR CONTRIBUTIONS

Tomoki Yaguchi: Data curation; formal analysis; investigation; visualization; writing-original draft; writing-review & editing. Shunsuke Kimura: Data curation; formal analysis; investigation; visualization; writing-original draft; writing-review & editing. Masahiro Sekiguchi: Data curation; formal analysis; investigation; visualization; writing-original draft; writing-review & editing. Yasuo Kubota: Investigation. Masafumi Seki: Data curation; formal analysis; investigation; visualization; writing-original draft. Kenichi Yoshida: Investigation. Yuichi

### TABLE 1

| Clone | Gene | Amino-acid change | Variant allele frequency (%) |
|-------|------|-------------------|-------------------------------|
|       |      |                   | D   | R2  | R3  | R4  | LN | Liver |
| 1     | BCOR | L1690_1721Ddup    | 42  | 32  | 34  | 40  | 46 | 38    |
| 2     | FLT1 | M148I             |     |     |     | 33  | 39 | 39    |
| 3     | JAK2 | L892V             |     |     |     | 37  | 41 | 50    |
| 4     | MYH7 | K542R             |     |     |     | 31  | 49 | 31    |
| 5     | PRKRIR | E158Q           |     | 34  |     |     |     |       |
| 6     | KMT2D | D3165fs          |     |     | 11  |     |     |       |
| 7     | TSHR | P3fs             |     |     | 8   |     |     |       |
| 8     | CDK6 | K93Q              |     |     | 5   |     |     |       |
| 9     | IL6ST | T178A           |     |     |     | 6   |     |       |
| 10    | NF1 | V2046L           |     |     | 7   |     |     |       |
| 11    | KDM6A | Q230L          |     |     |     |     | 23  |       |
| 12    | OBCN | R8618Q           |     |     |     |     | 13  |       |
| 13    | SYNE1 | A7572E          |     |     |     |     | 18  |       |

Abbreviations: D, initial diagnosis; dup, duplication; fs, frameshift.
DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

ETHICAL STATEMENT
The present study was approved by the Ethics Committee of the University of Tokyo. Informed consent for publication was obtained from the patient’s parents.

REFERENCES

1. Argani P, Perlman EJ, Breslow NE, et al. Clear cell sarcoma of the kidney: a review of 351 cases from the National Wilms Tumor Study Group Pathology Center. Am J Surg Pathol. 2000;24(1):4-18.

2. Furtwängler R, Gooskens SL, van Tinteren H, et al. Clear cell sarcomas of the kidney registered on International Society of Pediatric Oncology (SIOP) 93-01 and SIOP 2001 protocols: a report of the SIOP renal tumour study group. Eur J Cancer. 2013;49:3497-3506.

3. Gooskens SL, Furtwängler R, Vujanic GM, et al. Clear cell sarcoma of the kidney: a review. Eur J Cancer. 2012;48:2219-2226.

4. Seibel NL, Li S, Breslow NE, et al. Effect of duration of treatment on treatment outcome for patients with clear-cell sarcoma of the kidney: a report from the National Wilms’ tumor study group. J Clin Oncol. 2004;22(3):468-473.

5. Seibel NL, Chi Y-Y, Perlman EJ, et al. Impact of cyclophosphamide and etoposide on outcome of clear cell sarcoma of the kidney treated on the National Wilms Tumor Study-5 (NWTS-5). Pediatr Blood Cancer. 2019;66(1):e27450.

6. Gooskens SL, Furtwängler R, Spreafico F, et al. Treatment and outcome of patients with relapsed clear cell sarcoma of the kidney: a combined SIOP and AIEOP study. Br J Cancer. 2014;111:227-233.

7. Ueno-Yokohata H, Okita H, Nakasato K, et al. Consistent in-frame internal tandem duplications of BCOR characterize clear cell sarcoma of the kidney. Nat Genet. 2015;47(8):861-863.

8. Roy A, Kumar V, Zorman B, et al. Recurrent internal tandem duplications of BCOR in clear cell sarcoma of the kidney. Nat Commun. 2015;6:8891.

9. O'Meara E, Stack D, Lee C-H, et al. Characterization of the chromosomal translocation t(10;17)(q22;p13) in clear cell sarcoma of kidney. J Pathol. 2012;227(1):72-80.

10. Fehr A, Hansson MC, Kindblom L-G, et al. YWHAE-FAM22 gene fusion in clear cell sarcoma of the kidney. J Pathol. 2012;227:5-7.

11. Wong MK, Ng CCY, Kuick CH, et al. Clear cell sarcomas of the kidney are characterised by BCOR gene abnormalities, including exon 15 internal tandem duplications and BCOR–CCNB3 gene fusion. Histopathology. 2018;72:320-329.

12. Sekiguchi M, Seki M, Kawai T, et al. Integrated multiomics analysis of hepatoblastoma unravels its heterogeneity and provides novel druggable targets. NPJ Precis Oncol. 2020;4:20.

13. Shiraiishi Y, Sato Y, Chiba K, et al. An empirical Bayesian framework for somatic mutation detection from cancer genome sequencing data. Nucleic Acids Res. 2013;41:e89.

14. Dang HX, White BS, Foltz SM, et al. ClonEvol: clonal ordering and visualization in cancer sequencing. Ann Oncol. 2017;28(12):3076-3082.

15. Kimura S, Seki M, Yoshida K, et al. NOTCH1 pathway activating mutations and clonal evolution in pediatric T-cell acute lymphoblastic leukemia. Cancer Sci. 2019;110:784-794.

16. Vlad C, Gerrard M, Moertel C, et al. Treatment of recurrent clear cell sarcoma of the kidney with brain metastasis. Pediatr Blood Cancer. 2008;50(2):246-249.

17. Nishiyama H, Gill JH, Pitt E, Kennedy W, Knowles MA. Negative regulation of G1/S transition by the candidate bladder tumour suppressor gene DBCCR1. Oncogene. 2001;20:2956-2964.

18. Lopez-Beltran A, Alvarez-Kindelan J, Luque RJ, et al. Loss of heterozygosity at 9q32–33 (DBC1 locus) in primary non-invasive papillary urothelial neoplasm of low malignant potential and low-grade urothelial carcinoma of the bladder and their associated normal urothelium. J Pathol. 2008;215:263-272.

SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of this article.

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