Intraosseous inflammatory myofibroblastic tumor of the mandible with a novel ATIC-ALK fusion mutation: a case report

Yoko Tateishi1*, Koji Okudela1, Shigeo Kawai2, Takehisa Suzuki1, Shigeaki Umeda1, Mai Matsumura1, Mitomu Kioi3 and Kenichi Ohashi1

Abstract

Background: Inflammatory myofibroblastic tumor (IMT) is a rare low-grade malignant neoplasm with a predilection for children and young adults, and typically arises in the lung, abdominopelvic region, and retroperitoneum. IMTs in the maxillofacial region are extremely rare. Approximately 50% of IMT harbor rearrangements of the anaplastic lymphoma kinase (ALK) gene at 2p23 with various fusion partners.

Case presentation: We herein report a case of intraosseous IMT of the mandible with a novel ATIC-ALK fusion. Tooth 43 did not erupt after the loss of tooth 83 in an 11-year-old girl with no previous history of trauma. Panoramic tomography showed a unilocular radiolucent lesion in the right anterior mandible resorbing the root of tooth 42 and the medial side of the root of tooth 44. Computed tomography revealed a well-circumscribed 3-cm osteolytic lesion of the right anterior mandible eroding the buccal cortical plate. The entire lesion was curetted out. A histopathological examination revealed the proliferation of plump spindle cells with a storiform architecture and lymphocytes scattered around spindle cells. The spindle cells showed diffuse cytoplasmic staining for ALK by immunohistochemistry. A fluorescence in situ hybridization analysis revealed the translocation of a part of the ALK gene locus at chromosome 2p23. A rapid amplification of cDNA ends analysis confirmed the rearrangement of ALK and identified ATIC as a partner of this ALK fusion mutant.

Conclusion: To the best of our knowledge, this is the first case of intraosseous IMT of the mandible with a novel ATIC-ALK fusion. We also herein reviewed similar tumors reported in the literature.

Keywords: Inflammatory myofibroblastic tumor, Mandible, Anaplastic lymphoma kinase gene, ATIC-ALK fusion, Fluorescence in situ hybridization, 5′ rapid amplification of cDNA ends, RT-PCR

IMTs are mesenchymal neoplasms of intermediate malignant potential and histologically characterized by the proliferation of fibroblasts and myofibroblasts admixed with lymphocytes, plasma cells, eosinophils, and histiocytes [1]. The histological features of IMTs are often diagnostic, particularly through anaplastic lymphoma kinase (ALK) staining. The ALK gene has been implicated in the pathogenesis of IMTs, which supports the neoplastic origin of these tumors. Approximately 50% of IMTs harbor rearrangements in the ALK gene at 2p23 [4]. However, to the best of our knowledge, an ALK fusion mutation has not yet been demonstrated in IMTs in the jawbone.

We herein report a case of an intraosseous IMT in the mandible including a novel ATIC-ALK fusion mutation.
Case presentation
Clinical features
An 11-year-old girl presented with the delayed eruption of tooth 43 (two digit tooth notation system of the FDI) and mental nerve palsy. She had no history of trauma. An intraoral examination revealed that tooth 43 had not erupted after the loss of tooth 83 and the gingiva and alveolar mucosa of the right mandible were normal. Panoramic tomography showed a unilocular radiolucent lesion in the right anterior mandible resorbing the root of tooth 42 and the medial side of the root of tooth 44 (Fig. 1a). Tooth 43 was not impacted in the mandibular bone. Computed tomography showed a well-circumscribed non-expansile 3-cm osteolytic lesion in the right anterior mandible that eroded the buccal cortical plate and resorbed the roots of tooth 42 and tooth 44 (arrows) (Fig. 1b). Root resorption and cortical plate erosion suggested a locally aggressive tumor. The tumor lesion was entirely curretted out. No recurrence was detected in 18 months of follow-up.

Histopathological features and differential diagnosis
A histopathological investigation revealed plump spindle cell proliferation with a storiform architecture and lymphocytes scattered around the spindle cells (Fig. 2a, b). The background stroma included a focal myxoid matrix. These spindle cells had ovoid nuclei with a pale eosinophilic cytoplasm, suggesting a myofibroblastic, myoepithelial, or fibroblastic lineage. Small islands of an odontogenic epithelium were detected. No mitosis or necrosis was observed. The histomorphological findings suggested that spindle cells were of a myofibroblastic, myoepithelial, or fibroblastic lineage. Small islands of an odontogenic epithelium suggested that this tumor was an

Fig. 1 a Panoramic tomography showed the resorption of the root of tooth 42 and medial side of the root of tooth 44 (arrow heads). b Computed tomography revealed a well-circumscribed 3-cm osteolytic lesion in the right mandible that eroded the buccal cortical plate and resorbed the roots of tooth 42 and tooth 44 (arrows)

Fig. 2 Hematoxylin and eosin staining (a, b) and immunohistochemical staining (c, d). a The proliferation of plump spindle cells with a storiform architecture with scattered lymphocytes. Epithelioid components with an appearance suggestive of odontogenic epithelioid cells were detected. b Spindle cells contained ovoid nuclei with a pale eosinophilic cytoplasm. c Spindle cells were positive for SMA. d Spindle cells were positive for ALK
odontogenic tumor. High-grade malignant tumors were excluded because neither mitosis nor necrosis was observed. Differential diagnoses included myofibroma, myoepithelioma, an odontogenic tumor, solitary fibrous tumor, giant cell granuloma, and IMT.

**Immunohistochemical findings**

Immunohistochemical staining revealed that spindle cells were positive for α-smooth muscle actin (SMA) (Fig. 2c), ALK (Fig. 2d), CD10, and glial fibrillary acidic protein (GFAP), and negative for cytokeratin AE1/AE3, S100 proteins, and p63. The Ki67 labeling index was 3%.

**Molecular pathological findings**

**FISH**

A fluorescence in situ hybridization (FISH) (ALK) analysis was performed on a formalin-fixed paraffin-embedded (FFPE) tissue block. Rearrangements in the ALK gene at chromosome band 2p23 were detected by FISH utilizing a Vysis ALK break apart probe (Abbott Molecular). The FISH analysis revealed the translocation of a part of the ALK gene locus (Fig. 3a).

**RACE**

Total RNA was extracted from frozen tissue using an RNase Mini kit (Qiagen) according to the manufacturer’s instructions. In order to obtain cDNA fragments corresponding to a novel ALK fusion gene, a rapid amplification of cDNA ends (RACE) analysis was performed using the 5′-full RACE core set (TaKaRa Bio) according to the manufacturer’s instructions. First-strand cDNA synthesis was performed with a 5′-end-phosphorylated RT primer (5′-ATCTGGGCCCTTGTATTTATCCT). The primer set used for the first round of PCR was the A1 primer (5′-ACTTCCTGGTTGCTTTTGCGGGTTAT) and S1 primer (5′-AGAAGGAGCCACACAGAAGGGGTAA). The primer set used for the second round of PCR was the S2 primer (5′-CTCCTTCACAAACCAGAGACCAA) and A2 primer (5′-TTCAGGCAGGTCTTTCACAG). The second round PCR product covering the breakpoint was sub-cloned into a pT7Blue plasmid vector (TaKaRa Bio). The DNA sequence was analyzed using universal primers (M13M4 primer or T7 promoter primer) and the Big-dye terminator ver3.1 kit (Applied Biosystems). A RACE analysis confirmed a rearrangement in ALK and identified ATIC (NM_004044) as a partner of this ALK fusion mutant (Fig. 3b).

**RT-PCR**

First-strand cDNA was synthesized from total RNA using the SuperScript First-Strand Synthesis System (Invitrogen) according to the manufacturer’s instructions. RT-PCR was used to confirm the presence of the ATIC-ALK fusion gene in this case. The primer set used was as follows: forward (5′-CCAAGGCTCCATCACAGTGTTACC) and reverse (5′-GAGGTCTTTGCGCAAG GACGTAGT). RT-PCR was used to confirm the presence of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) in this case. The primer set used was as follows: forward (5′-TGGAGTCCACTGCGGCTTTC) and reverse (5′-ATGACGACATGGGGGCATCAG). A RT-PCR analysis confirmed the ATIC-ALK fusion in tumor cells (Fig. 3c).

Based on these results, the final diagnosis was IMT.

**Discussion**

We herein presented the first case of IMT with an ATIC-ALK fusion mutation primarily in the maxillofacial region. IMTs are rare mesenchymal tumors. Different terms have been applied to these lesions, namely, xanthogranuloma, plasma cell granuloma, inflammatory...
pseudotumor (IPT), and inflammatory myofibroblastic sarcoma [1, 6, 7]. The diverse nomenclature is descriptive and reflects the uncertainty regarding the true biological nature of these lesions. In 2013, the World Health Organization classification of soft tissue and bone defined IMT as a distinctive neoplasm composed of myofibroblastic and fibroblastic spindle cells accompanied by an inflammatory infiltrate of plasma cells, lymphocytes, and/or eosinophils [1]. The expression of ALK proteins is detected by anti-ALK immunohistochemistry in approximately 50% of IMT cases. ALK protein expression has been shown to reliably correlate with ALK gene rearrangements [8]. These lines of evidence support ALK-positive IMT being a distinct neoplastic entity. ALK immunostaining is an aid in the pathological diagnosis of IMTs, as well as in the differentiation of IMTs from other spindle cell neoplasms that fall within the broad category of IPTs. We reviewed 12 cases of intraosseous tumors of the mandible with similar histological features in the literature (Table 1) [7, 9–19]. Six cases were diagnosed as IPTs and 6 as IMTs in the mandible. ALK-positive IMT/IPT in the mandible was initially reported in 2005 [10] and only 4 cases including ours have been reported since then [10, 11, 19].

Thus, IMT may very rarely occur in the mandible. Otherwise, IMT may often be misdiagnosed because difficulties are associated with its histological differentiation from other spindle cell neoplasms and reactive lesions. Differential diagnoses include posttraumatic spindle cell nodules, giant cell granuloma, myofibroma, myoepithelioma, ameloblastic fibroma, odontogenic myxofibroma, IMT, a solitary fibrous tumor, low-grade myofibroblastic sarcoma, myofibrosarcoma, leiomyosarcoma, and spindle cell carcinoma. The immunohistochemical expression of the ALK protein is a very important hallmark for differentiating IMT from those described above. Therefore, if spindle cell lesions are detected in the jawbone, IMT must be included in the differential diagnosis and an immunohistochemical examination of ALK protein expression is needed.

### Table 1 Clinicopathological features of IMTs in the mandible described in the present and previous reports

| No. | Authors (yr.) | Age (yr.) | Sex | Diagnosis | ALK-IHC | FISH | RT-PCR |
|-----|---------------|-----------|-----|-----------|---------|------|--------|
| 1   | Zegarelli (1974) [7] | 56 | F | Granuloma (?IPT) | NA | NA | NA |
| 2   | Inui (1993) [9] | 63 | M | IPT | NA | NA | NA |
| 3   | Brooks (2005) [10] | 82 | F | IMT | positive | NA | NA |
| 4   | Poh (2005) [11] | 42 | F | IMT | positive | NA | NA |
| 5   | Oh (2008) [12] | 20 | F | IPT | NA | NA | NA |
| 6   | Johann (2008) [13] | 33 | M | IPT | NA | NA | NA |
| 7   | Satomi (2010) [14] | 14 | F | IMT | negative | NA | NA |
| 8   | Date (2010) [15] | 70 | M | IPT | negative | NA | NA |
| 9   | Sasagawa (2011) [16] | 60 | M | IMT | NA | NA | NA |
| 10  | Gawande (2012) [17] | 20 | M | IPT | NA | NA | NA |
| 11  | Sah (2013) [18] | 30 | M | IMT | negative | NA | NA |
| 12  | Stringer (2014) [19] | 16 | M | IMT | positive | NA | NA |
| 13  | Present case | 11 | F | IMT | positive | positive | ATIC-ALK |

ALK anaplastic lymphoma kinase, IHC immunohistochemistry, FISH Fluorescence in situ hybridization, F female, M male, IPT Inflammatory pseudotumor, IMT Inflammatory myofibroblastic tumor, NA data not available

Fig. 4 The hypothetical schema of the gene fusion between ALK and ATIC. ATIC-ALK fusion is generated by the pericentric inversion of chromosome 2 at the breakpoints of p23 and q35 (inv(2)(p23q35)), in which the ALK and ATIC genes are localized.
The present tumor was confirmed to have the ATIC-ALK fusion mutation. Different ALK fusion partners, including TPM3, TPM4, CRTC, CARS, RANBP2, ATIC, SEC31L1, PPFIBP1, and DCTN1, in which TPM3 is the most common partner, have been identified in IMTs [5, 20–26]. ATIC-ALK fusion mutations are very rare, comprising approximately 1% of ALK-fusion mutations in anaplastic large cell lymphomas (ALCL) [27]. The hypothetical schema of the gene fusion between ATIC and ALK is shown in Fig. 4 [28]. To the best of our knowledge, we are the first to describe IMT with the ATIC-ALK fusion mutation and one case of IMT arising from the hyoid bone in a neonate with the ATIC-ALK fusion have been reported in the literature [23, 29]. Thus, the present case is the first IMT with the ATIC-ALK fusion mutation primarily in the mandible.

The management of IMTs needs to entail complete surgical resection, including postoperative reassessments for at least 10 years [30]. Although the number of IMTs in the jawbone is limited, they appear to exhibit a favorable clinical course. However, it is important to emphasize that IMTs occasionally behave in an aggressive manner. Poh et al. reported the local recurrence of an intraosseous IMT 14 months after initial surgery [11]. Sasagawa et al. described a case of multiple IMTs involving the cranium, mandible, ischium, and calcaneus [16]. The ALK inhibitor, crizotinib, was recently demonstrated to be clinically beneficial for IMTs with ALK fusion mutations [26, 31–33].

Conclusion

IMTs are rare lesions in the mandible. The general consensus is that IMTs encompass a group of heterogeneous lesions with various biological behaviors. However, a proportion of cases are genuine neoplasms with the involvement of ALK fusion mutations. The recognition of this diagnostic entity is important for selecting appropriate treatments. To the best of our knowledge, we are the first to describe IMT with the ATIC-ALK fusion mutation in the mandible.

Abbreviations

IMT: Inflammatory myofibroblastic tumor; ALK: Anaplastic lymphoma kinase; FISH: Fluorescence in situ hybridization; RACE: Rapid amplification of cDNA ends; RT-PCR: Reverse transcription polymerase Chain reaction; IFT: Inflammatory pseudotumor

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Availability of data and materials

The dataset supporting the conclusions of this article is included within the article.

Authors’ contributions

YT and KO carried out the molecular genetic studies and drafted the manuscript. SK provided histological and immunohistochemical evaluation. TS, SU and KM carried out the molecular genetic studies. MK attended the patient and arranged the clinical examination. KO conceived of the study, and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

Written informed consent was obtained from the patient’s family for the publication of this case report and any accompanying images.

Ethics approval and consent to participate

Not applicable for Case Report.

Author details

1Department of Pathology, Yokohama City University Graduate School of Medicine, Yokohama, Japan. 2Department of Pathology, Japanese Red-Cross Musashino Hospital, Tokyo, Japan. 3Department of Oral and Maxillofacial Surgery, Yokohama City University Graduate School of Medicine, Yokohama, Japan.

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