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Diagnostic value of *H3F3A* mutations in giant cell tumour of bone compared to osteoclast-rich mimics

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Abstract

Driver mutations in the two histone 3.3 (*H3.3*) genes, *H3F3A* and *H3F3B*, were recently identified by whole genome sequencing in 95% of chondroblastoma (CB) and by targeted gene sequencing in 92% of giant cell tumour of bone (GCT). Given the high prevalence of these driver mutations, it may be possible to utilise these alterations as diagnostic adjuncts in clinical practice. Here, we explored the spectrum of *H3.3* mutations in a wide range and large number of bone tumours ($n = 412$) to determine if these alterations could be used to distinguish GCT from other osteoclast-rich tumours such as aneurysmal bone cyst, nonossifying fibroma, giant cell granuloma, and osteoclast-rich malignant bone tumours and others. In addition, we explored the driver landscape of GCT through whole genome, exome and targeted sequencing (14 gene panel). We found that *H3.3* mutations, namely mutations of glycine 34 in *H3F3A*, occur in 96% of GCT. We did not find additional driver mutations in GCT, including mutations in *IDH1*, *IDH2*, *USP6*, *TP53*. The genomes of GCT exhibited few somatic mutations, akin to the picture seen in CB. Overall our observations suggest that the presence of *H3F3A* p.Gly34 mutations does not entirely exclude malignancy in osteoclast-rich tumours. However, *H3F3A* p.Gly34 mutations appear to be an almost essential feature of GCT that will aid pathological evaluation of bone tumours, especially when confronted with small needle core biopsies. In the absence of *H3F3A* p.Gly34 mutations, a diagnosis of GCT should be made with caution.

Keywords: *H3F3A*; *H3F3B*; giant cell tumour of bone; malignant giant cell tumour of bone; giant cell granuloma; solid variant of aneurysmal bone cyst; *USP6*

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Introduction

Giant cell tumours of bone (GCT) account for approximately 5% of all primary bone tumours and generally affect the epiphysis of long bones in skeletally mature individuals. They are classified as locally aggressive and rarely metastasising neoplasms [1]. When GCT metastasise, they spread to the lungs but notably the morphology remains similar to the primary lesion and the disease is generally controlled by resection alone. Such an event is referred to as 'benign metastasing' disease. Although the majority of GCTs presents with a characteristic combination of clinical, radiological and histological findings, some remain diagnostically challenging, particularly on small needle core biopsies [1]. Due to varying histological appearances within and between individual GCT and due to morphologic overlap with other bone tumours, the differential diagnoses include osteoclast-rich osteosarcomas which often represent telangiectatic osteosarcoma, and high grade sarcoma arising within a conventional GCT (malignant GCT), along with several benign mimics comprising aneurysmal bone cyst (ABC), chondroblastoma (CB), nonossifying fibroma (NOF) and giant cell granuloma (GCG) occurring exclusively in the jaws and in the small tubular bones of the hand and feet [2,3]. Less likely, GCT can also be mistaken for osteoid osteoma/osteoblastoma and tenosynovial giant cell tumour.

Recently, we reported the whole genomes of 6 CB which harboured p.Lys36Met (p.K36M) mutations in the paralogous histone 3.3 genes (*H3.3*), *H3F3A* (1/6) and *H3F3B* (5/6). The finding was confirmed in an extended targeted sequencing analysis detecting these mutations in 73/77 CB. Due to similarities in morphology and site of presentation, we also screened a series of GCT for *H3.3* mutations. We found *H3.3* mutations in 92% of cases (49/53), exclusively in the *H3F3A* gene which all occurred at glycine 34. *H3F3A* p.Gly34Trp (p.G34W) alterations were found in all but one case of GCT which harboured a p.G34L mutation. We additionally identified distinct *H3.3* mutations in 2/103 osteosarcomas (one p.G34R in *H3F3A*, and one p.G34R mutation in *H3F3B*) and in 1/75 chondrosarcomas (*H3F3A* p.K36M) [2]. Since then, others have reported a *H3F3A* mutation in another osteosarcoma (1/10, p.G34W) [3].

In addition to *H3F3A* and *H3F3B* mutations, other somatic mutations may potentially be of diagnostic value in the clinic for classifying osteoclast-rich lesions of bone. Such mutations include rearrangements involving *USP6* which forms a fusion gene with a number of partners including *CDH11*, *ZNF9*,

COL1A1, *TRAP150* and *OMD*. These are found in approximately 75% of primary ABC [4–6] but have not been found in a screen of 9 GCT [4,5]. In addition, fusions involving macrophage colony-stimulating factor 1 have been described in tenosynovial giant cell tumours. Unfortunately this is of limited diagnostic value as it is only detectable in a minority of tumour cells [7]. Furthermore, *IDH2* mutations, which also affect central chondrosarcoma [8], have recently been reported to occur in 80% of GCT (16/20) [9].

In this study, we sequenced the whole genomes of five and whole exome of one GCT to search for driver mutations other than H3.3 mutations. In addition, we sought to define the specificity of p.G34W alterations for GCT by screening 91 GCT as well as a series of 321 osteoclast-rich lesions including ABC, CB, NOF, GCG, osteoblastoma/osteoid osteoma, TGCT, in addition to malignant osteoclast-rich tumours of bone.

Materials and methods

Patient samples

Formalin-fixed paraffin-embedded cases were retrieved from the diagnostic pathology archives at the Royal National Orthopaedic Hospital (RNOH) and from the Basel Bone Tumour Reference Centre (BBTRC). All samples were reviewed by specialist bone tumour pathologists at RNOH (AMF, MFA, RT) and BBTRC (DB) and diagnosed according to the current WHO classifications [10,11] (Supporting Information Tables 1 and 2). Individual cases were correlated with clinical and radiological information where available. Tissue sample types are provided in Supporting Information Table 3. In cases in which the histology and mutational analysis did not match, the morphology was re-evaluated independently by all members of the RNOH pathology team and a consensus diagnosis was subsequently reached by all pathologists.

Ethical approval was obtained from the Cambridgeshire 2 Research Ethics Service (reference 09/H0308/165), the UCL Biobank for Health and Disease ethics committee (covered by the Human Tissue Authority licence 12055: project EC17.1) and from the Ethikkommission beider Basel (reference 274/12).

Whole genome and exome sequencing

DNA from GCT and matching non-neoplastic tissue, derived from 6 individuals, was subjected to whole genome ($n = 5$) or exome ($n = 1$) sequencing as described previously [12], using Illumina HiSeq 2000

Table 1. Overview of giant cell tumours of bone genomes and exomes

	PD21292a	PD21294a	PD21295a	PD21296a	PD7524a	PD9999a
Genome/exome	Genome	Genome	Genome	Genome	Genome	Exome
Tumour coverage	51.26	41.42	56.22	48.48	39.04	~60% covered by at least 30×
Normal coverage	25.93	22.56	23.28	26.1	34.26	~60% covered by at least 30×
Copy number source	Sequencing reads	Sequencing reads	Sequencing reads	Sequencing reads	SNP6 and sequencing reads	n.a.
Structural changes	0	0	0	0	LOH 5q and 8p	n.a.
Substitutions	479	977	739	700	815	7
Precision*	0.90	0.90	0.93	0.90	0.81	All reviewed
Coding	5	6	9	11	12	7
Non-synonymous	4	4	7	7	8	7
Truncating	0	1	0	1	2	0
Indels	54	79	102	46	45	0
Precision*	0.76	0.84	0.61	0.91	0.93	n.a.
Coding	0	0	0	0	0	0
In frame	0	0	0	0	0	0
Out of frame	0	0	0	0	0	0
Driver mutation	H3F3A G34W	H3F3A G34W**	H3F3A G34W	H3F3A G34W	H3F3A G34W	H3F3A G34W

*Based on review of 100 (or less, where fewer mutations were called) randomly selected variants.

**Manually called.

n.a.—not applicable.

analysers (Illumina Inc, CA). Reads were aligned to the reference human genome (NCBI37) by burrows wheeler aligner (BWA) on default settings [13]. Unmapped reads and polymerase chain reaction-derived duplicates were excluded. Relevant coverage metrics are summarised in Table 1.

Targeted next generation sequencing with Ion Personal Genome Machine® (PGM™)

A pooled barcoded amplicon-tagged library generated using Fluidigm® Access Array™ (Fluidigm Europe B.V, Amsterdam, Netherlands) was diluted for direct

input into the emulsion polymerase chain reaction (PCR) with Ion Sphere™ particles with Ion PGM™ Template OT2 200 Kit (Ion OneTouch™ system) following the manufacturer’s instructions (Life Technologies Ltd, Paisley, Scotland). The pooled samples were sequenced using 316 chip with Ion PGM™ 200 Sequencing kit on the Ion Torrent PGM™ sequencer following manufacturer instructions (Life Technologies). Targeted hotspot mutations as reported in COSMIC in 14 genes, including *EGFR*, *KRAS*, *NRAS*, *BRAF*, *IDH1*, *IDH2*, *KIT*, *GNAS1*, *CTNNB1*, *PTEN*, *PDGFRA*, *H3F3A*, *H3F3B*, *TP53* (covering in total 50 amplicons) were analysed (Supporting

Table 2. Results of the targeted genetic analysis for H3F3A and H3F3B and USP6 rearrangement

Tumour type	Mutation characteristic of tumour									*Revised diagnosis (n)
	Original Dx (n)	GCT				CB		ABC	WT	
		G34W/A (n)	G34L/A (n)	G34R/A (n)	G34M/A (n)	K36/A (n)	K36/B (n)	USP6 (n)		
GCT	91	83	0	1	1	0	0	2	4	89
GCT of small bones	25	14	2	0	0	0	0	3	6	21
CB	7	0	0	0	0	5	2	0	0	7
ABC	44	2	0	0	0	0	1	32	9	41
NOF	66	4	0	0	0	0	0	0	62	62
Tenosynovial giant cell Tumour	51	0	0	0	0	0	0	0	51	51
Osteoblastoma	21	0	0	0	0	0	0	0	21	21
Osteoid osteoma	9	0	0	0	0	0	0	0	9	9
Giant cell granuloma of jaw (solitary)	78	0	0	0	0	0	0	0	78	78
Adamantinoma	8	0	0	0	0	0	0	0	8	8
Osteofibrous dysplasia	6	0	0	0	0	0	0	0	6	6
Total no of mutations		103	2	1	1	5	3	37	252	

*Diagnosis revised following review of histology and imaging on discovery of the genetic alterations

Table 3. Mutation analysis results for the 10 malignant osteoclast cell-rich tumours

ID	Year of primary and relapses	Alive/DOD	Age of diagnosis	Gender	Site of primary tumour	Histopathological diagnosis	TP53	Histone variant	IDH1/2 alterations
S00033676	2012	ALIVE	33	M	Talus, left	Malignant osteoclast-rich tumour	WT	H3F3A/B WT	IDH1/2 WT
S00033682	2005	Lost in follow up	59	F	Fibula, right, proximal	Osteoclast-rich dedifferentiated chondrosarcoma	WT	H3F3A/B WT	IDH2 R132S
S00033684	1998, 2009	ALIVE	37	F	Toe, 1st, left, proximal	Malignant osteoclast-rich tumour with features of osteosarcoma	WT	H3F3A/B WT	IDH1/2 WT
S00033694	2006, 2008, 2010, 2011, 2013	DOD (11/2013)	52	F	Tibia, left, proximal	Malignant osteoclast-rich tumour with features of osteosarcoma	WT	H3F3A/B WT	IDH1/2 WT
S00039478	2006	Alive	11	M	Metatarsal, 3rd, 4th, 5th, left	Malignant osteoclast-rich tumour with features of osteosarcoma	WT	H3F3A/B WT	IDH1/2 WT
S00039480	2002, 2006, 2010	Lost in follow up	39	M	Femur, left, distal	Malignant osteoclast-rich tumour, multicentric	WT	H3F3A/B WT	IDH1/2 WT
S00039484	2004	DOD (10/2007)	74	M	Femur, left, distal	Malignant osteoclast-rich tumour with features of osteosarcoma	WT	H3F3A/B WT	IDH1/2 WT
S00039476	1998, 2005, 2009	Alive	19	F	Tibia, distal	Malignant osteoclast-rich tumour with features of osteosarcoma	WT	H3F3A G34W/A	IDH1/2 WT
S00038597	2010	DOD	40	F	Tibia, right, proximal	Dedifferentiated adamantinoma	WT	H3F3A G34W/A	IDH1/2 WT
S00033295	2013, 2014	Alive	50	M	Tibia, left, distal	Malignant osteoclast-rich tumour	WT	H3F3A/B WT	IDH1/2 WT

Information Table 4 for primer details and description of hot-spots targeted).

Reads were aligned to the reference genome hg19 and binary alignment map (BAM) files were generated using Ion torrent suite version 4.0.1. Reads were visualised using integrative genomics viewer (IGV) [14] with the appropriate browser extensible data (BED) files for all 14 genes.

Results

Whole genome/exome sequencing of GCT

To perform a genome-wide search for driver mutations in GCT, DNA from 5 tumours and blood from

the same patients, were studied by whole genome sequencing, and one tumour was subjected to whole exome sequencing. Overall, few somatic changes were seen. In the five genomes, a median of 739 substitutions (range 479–977) and 54 indels (range 45–102) were present (Supporting Information Table 5). Including the additional tumour studied at whole exome level, overall 50 coding substitutions, 37 of which were nonsynonymous and no coding indels were found. Other than *H3F3A* p.G34W mutations, no driver event was discernible amongst nonsynonymous mutations. No rearrangements or copy number changes were identified, bar two regions of loss of heterozygosity in one of the five genomes (PD7524a) (Supporting Information Figure 1).

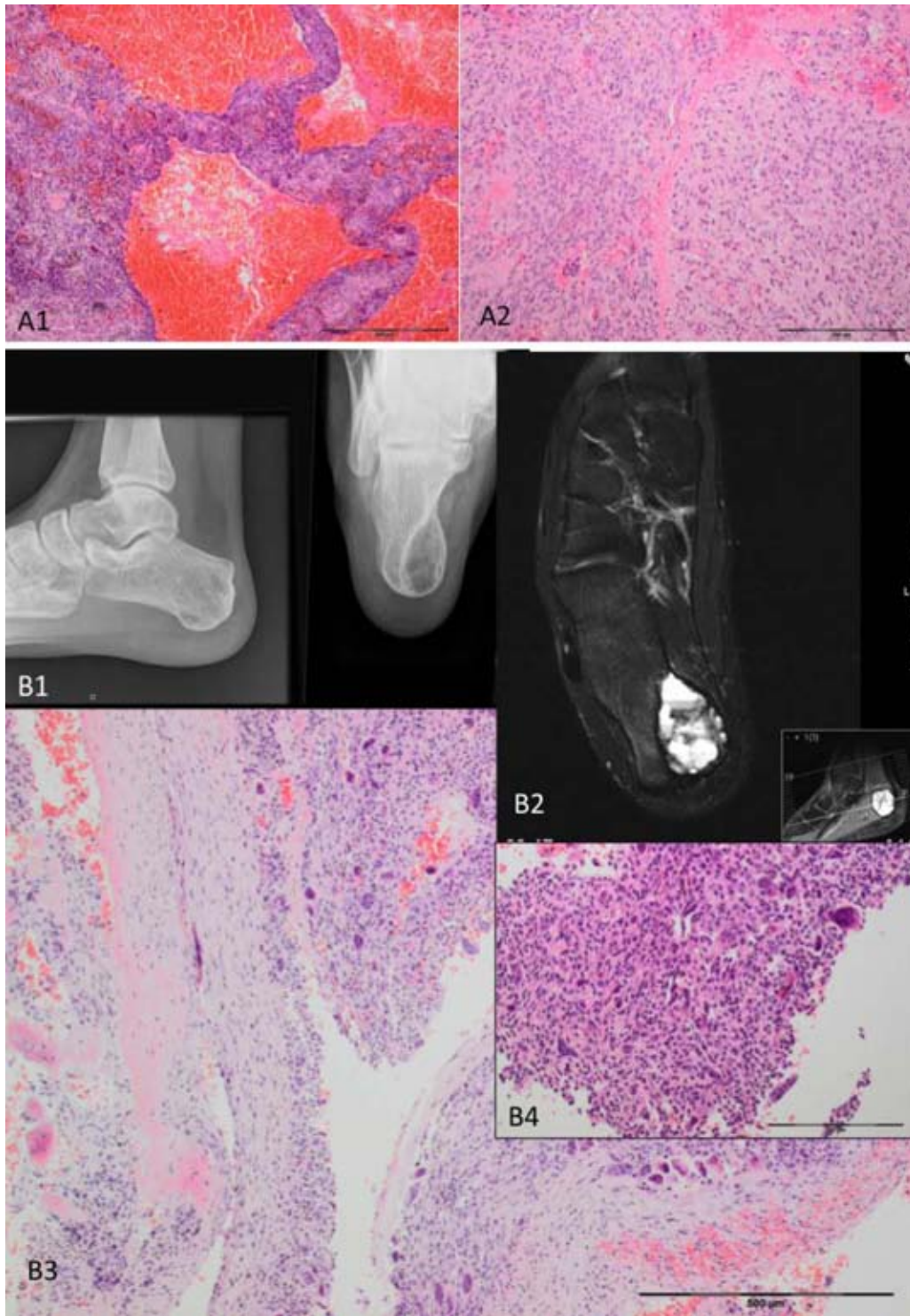


Figure 1. A sacral tumour (S00036624) (A) and a calcaneal tumour (S00036606) (B) originally diagnosed as ABCs. A sacral tumour in an 18-year-old female originally diagnosed with an ABC. The tumour shows blood-filled vascular spaces contained by septa (A1), and scattered solid areas (A2), both of which contain numerous osteoclasts: a *USP6* gene rearrangement is not detected. The tumour harboured a *H3F3A* p.G34W mutation after which the lesion was then reclassified as a giant cell tumour (GCT) with ABC change. A calcaneal tumour in a 22-year-old male originally diagnosed with an ABC. The X-ray reveals a well-defined lucency (B1), and the MRI shows fluid–fluid levels (B2). The histology shows an osteoclast-rich predominantly cystic lesion (B3, B4). A *H3F3B* p.K36M mutation was detected. The histology was reviewed and small areas with features of CB (B4) were detected. The tumour was reclassified as CB with extensive ABC change.

Targeted genetic analysis

Giant cell tumour of bone. A histological diagnosis of GCT was favoured in 91 osteoclast-rich tumours: 83 were found to harbour a *H3F3A* p.G34W alteration and alterations represented by p.G34R and p.G34M were detected in two tumours (Table 2). No mutations were detected in the *H3F3B* gene. The two GCTs (femur S00033722, ulna S00033686) which metastasised to the lung harboured a *H3F3A* p.G34W mutation. Morphologically, both primary tumours and the metastases showed the typical features of a conventional GCT. Both patients are alive and well and the tumours were classified as 'benign metastasising' GCT [15].

Fluorescence in situ hybridisation (FISH) revealed *USP6* break-apart signals in 2/6 remaining wild type (WT) GCT, both of which were located in the spine (Table 2). In these 2 cases, the histology was re-evaluated and considered compatible with ABC and the diagnosis revised accordingly. Ten GCT with a *H3F3A* p.G34W were also tested for *USP6* rearrangements and all were negative. Consequently, 85/89 (96%) GCT showed mutations in *H3F3A* genes (Table 2).

Sixty GCT were screened for hotspot mutations in 12 genes in addition to *H3F3A* and *H3F3B* using the Ion Torrent PGM. No mutations, specifically no *TP53*, *IDH1* and *IDH2* alterations, were identified (Supporting Information Table 2).

None of the 89 GCT presented in patients with immature skeleton, although 2 female patients presented at the age of 15 and one at 17.

Chondroblastoma. All seven CB analysed by targeted sequencing harboured a p.K36M mutation, five of which were in the *H3F3A* and two in the *H3F3B* gene (Table 2). Seven CB were analysed for *USP6* rearrangements, including 4/77 cases that were WT for *H3F3A* and *H3F3B* in our previous study (Table 1), and the 3/7 cases from this study in which a p.K36M alteration was identified (Table 2). None were detected. No mutations in the hotspots in the Ion Torrent PGM 14 gene panel assay were detected in 19 CB harbouring p.K36M: these included 18 cases from the previous study (Supporting Information Table 1, 2).

Aneurysmal bone cyst. Of 44 tumours where the favoured initial histological diagnosis was ABC, 32 revealed a *USP6* rearrangement by FISH: 27 showing break-apart signals, with 5 showing deletions of the telomeric end (Figure 1, Table 2). One of the remaining cases (1/12) harboured a *H3F3B* p.K36M mutation which on histological review revealed a tiny fragment with features of a CB (Figure 1B). The diagnosis was, therefore, revised. Two additional

cases (2/12) revealed *H3F3A* p.G34W mutation and were considered to be compatible with GCT on histologic re-evaluation (Figure 1A). The remaining 9/12 cases were WT for both the *USP6* and *H3.3* alterations. The *USP6* alterations were mutually exclusive with *H3.3* mutations. Reclassification of ABC on the basis of the genetic finding in 3/44 lesions resulted in a diagnosis of 41 ABC, 32 (78%) of which demonstrated *USP6* rearrangements (Table 2).

Osteoclast-rich tumours of the small bones of hand and feet and patella. Twenty-eight osteoclast-rich tumours of small bones of the hand ($n = 13$), feet ($n = 14$) and patella ($n = 1$) were studied, including three cases from our previous study (Supporting Information Table 1, 2). Sixteen tumours harboured a *H3F3A* mutation, 14 of which involved a p.G34W alteration. The other two alterations were represented by p.G34L mutations (Supporting Information Table 1, 2). One tumour in the talus harboured a *H3F3B* p.K36M substitution. On histological review a small focus with features of a CB was seen and the tumour was reclassified accordingly. *USP6* break-apart signals were detected in 3/11 remaining cases that were WT for *H3F3A* and *H3F3B*, and these were classified as ABC, leaving 8/28 (29%) cases (patella, bone of hand $\times 2$, bone of foot $\times 5$) without an alteration with which to classify the disease. The *USP6* rearrangement was mutually exclusive with *H3.3* mutations in these cases. Screening of the mutation hotspots on Ion Torrent PGM failed to reveal any additional mutations in the 24/28 cases.

Nonossifying fibroma, tenosynovial giant cell tumour, osteoblastoma, osteoid osteoma, and giant cell granuloma of the jaw. Details of these tumours are included in Supporting Information Table 2 and genetic alterations in Table 2: 4/66 previously diagnosed NOF harboured *H3F3A* p.G34W alterations. These were reclassified as GCT following review of histology and imaging. No *H3F3A* and *H3F3B* mutations or hotspot mutations were identified in the Ion Torrent screen. Fifteen GCG of the jaws analysed by FISH for *USP6* rearrangements did not reveal a break-apart signal or deletion of one the probes.

We screened 51 tenosynovial giant cell tumours, 21 osteoblastomas, 9 osteoid osteomas, and 78 GCG of the jaw, three cases of genetically proven cherubism and two cases of genetically proven Noonan syndrome. No p.G34 alterations in *H3F3A* or p.K36 alterations in the *H3F3B* were identified (Table 2).

H3F3A and H3F3B mutations in malignant osteoclast-rich tumours. Ten cases classified as malignant osteoclast-rich tumours were retrieved from the

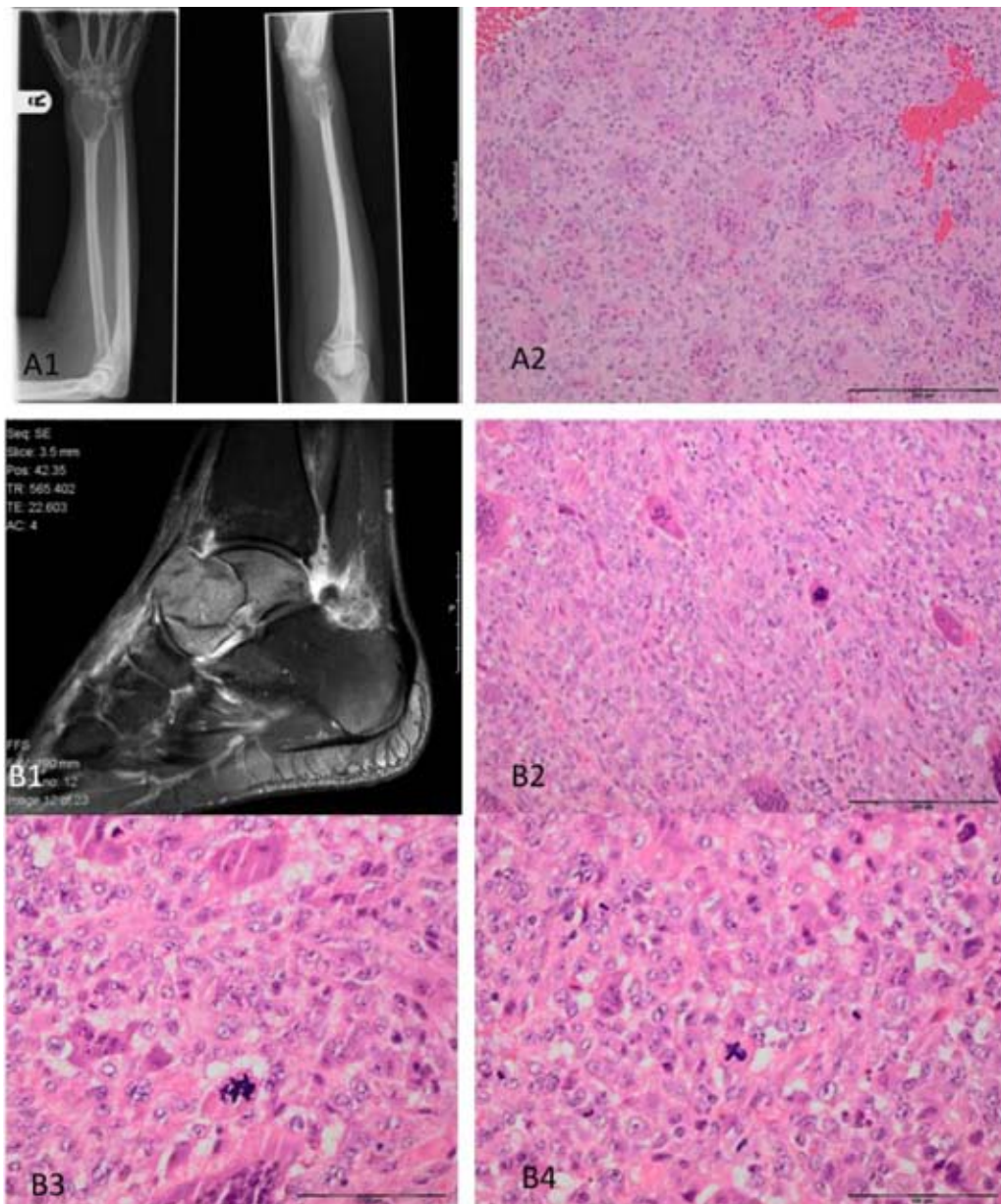


Figure 2. Photomicrographs and X-rays of a conventional (S00033682) (A), and malignant giant cell tumour (GCT) (S00030176) (B). A central lytic tumour of the distal radius (A1). The histology is typical for GCT in which the stromal cells show no significant cytological atypia (A2). An aggressive lytic lesion of the talus, which has broken through the cortex (B1). Microscopy shows features of a malignant osteoclast-rich tumour in which the stromal cells are enlarged with nuclear pleomorphism and numerous mitoses, including atypical forms (B3 and B4). Geographic-type tumour necrosis was also present (not shown).

RNOH archives, eight of which were wild type (WT) for *H3F3A* and *H3F3B* mutations (Table 3). On histological review, four cases revealed features typical of osteosarcoma in addition to osteoclast-rich areas. These cases were reclassified as osteosarcoma with osteoclast-rich areas. Of the four other WT cases, three were composed predominantly of areas of malignant GCT (Figure 2B). The remaining tumour

was a predominantly osteoclast-rich tumour which had been reported as being 'suspicious for malignancy' and revealed an *IDH2* p.R172S mutation. On review of the biopsy, a small fragment of mature neoplastic cartilaginous tissue in addition to the osteoclast-rich neoplasm was noted. The tumour was mitotically active (20 mitoses/high power fields). No cartilage was detected in the resection specimen

although an area showing a monotonous spindle cell overgrowth was present (Supporting Information Figure 2). This histological observation associated with the *IDH2* p.R172S mutation supported a diagnosis of a dedifferentiated chondrosarcoma in which the dedifferentiated component was an osteoclast-rich neoplasm. The tumour was reclassified accordingly.

Of the three patients whose tumours showed predominantly a malignant GCT component, one was sited in the talus and the patient is well having experienced no recurrence following amputation with 24 months of follow-up (Figure 2B). The second was a multicentric malignant GCT diagnosed in 2002. This patient developed brain and lung metastasis and died 9 years after diagnosis. The third tumour developed in the distal tibia and within 6 months of diagnosis and amputation, the patient developed recurrent disease at the amputation stump and metastatic disease in the pelvis.

One osteoclast-rich malignant tumour which harboured a *H3F3A* p.G34W mutation originated in the distal tibia and within a short interval presented with multifocal disease in the vertebrae, which revealed unequivocal malignant features. The patient was treated with resection and radiotherapy and is alive with disease 20 years after diagnosis. Neither tissue nor imaging from the original tumour was available for review.

The second osteoclast-rich malignant tumour which harboured a *H3F3A* p.G34W mutation was classified as an unusual, high grade osteoclast-rich tumour with adamantinoma-like features located in the right proximal tibia. Histological features typical of a conventional adamantinoma were present and this component merged with a high grade spindle cell sarcomatous area partly infiltrated with large numbers of osteoclasts. Sited in the subarticular space it was considered more in keeping with a GCT because adamantinomas generally occur in the diaphysis (Supporting Information Figure 2). No *TP53* mutations were detected. The tumour was resected *en bloc*: the patient developed multiple spinal metastases and died of disease 7 months later. The different histological areas were microdissected and all revealed the *H3F3A* p.G34W alteration (Table 3). *H3.3* mutations were not detected in six other cases of osteofibrous dysplasia and eight cases of adamantinoma (Table 2).

Overall our observations suggest that the presence of *H3F3A* p.G34W mutations does not entirely exclude malignancy in osteoclast-rich tumours. However, *H3F3A* p.G34W mutations appear to be an almost essential feature of GCT that will aid pathological evaluation of bone tumours, especially when confronted with small needle core biopsies (see

algorithm-based for diagnosis decision Supporting Information Figure 3).

Discussion

Whole genome/exome sequencing of 6 GCT showed remarkably few somatic changes, other than the occurrence of *H3F3A* mutations. In this way, GCT genomes showed similarities with previously reported CB genomes [2] which reflects overlapping features of the two tumour types, including the osteoclast-rich morphology, their site of occurrence and their biological behaviour. The notable difference between these tumours is the association of cartilaginous differentiation in CB with the *H3.3* K36M alteration, the majority of which occur in *H3F3B*, whereas 96% of the mutations in GCT occur in *H3F3A* involving p.G34. The finding of highly recurrent *H3.3* substitutions in the absence of other recurrent alterations in both primary and recurrent disease also implies that these mutations in both GCT and CB represent essential oncogenic drivers. It is noteworthy that in the two cases of 'benign' metastatic disease no driver mutation other than p.G34W alterations were found in the primary and distant lesions.

In our previous publication [2], we remarked on the exquisite specificity of individual *H3.3* mutations for different bone and brain tumour types. Here, we confirm that the vast majority (83/89) of GCT alterations occur in *H3F3A* and involve p.G34W. However, we also detected an overlap of specific *H3.3* mutations and tumour types in a minority of cases. We found that p.G34 substitutions in GCT involve on rare occasions *H3F3A* p.G34R which is commonly seen in paediatric glioblastoma. In addition, p.G34M was also detected in a single GCT, a substitution previously reported only once in diffuse intrinsic pontine glioma [16]. In contrast to the occurrence of p.G34R and p.G34M in aggressive brain tumours, in which *TP53* mutations are also seen [17], the 3 GCT with these alterations followed a typical clinical course and did not harbour *TP53* mutations, with 2/3 having at least 5 years follow-up (p.G34R—2009; p.G34M—2004). Finally, we also confirm that p.G34L mutations are rare and 2/3 occurred in small bones of the hands and feet. This mutation, like p.G34W, appears to be confined to bone tumours, having not been reported in primary brain tumours.

In contrast to the high prevalence of *H3.3* mutations in GCT and CB, these are rare in malignant primary bone tumours. Specifically, *H3.3* mutations have previously been reported to occur in 3/113

osteosarcoma, 1/15 clear cell chondrosarcoma and 1/75 central chondrosarcoma [2,3]. A novel finding in this study is the presence of a *H3F3A* p.G34W alteration in an unusual high grade osteoclast-rich tumour with adamantinoma-like features thereby providing further evidence to link an osteoclast-rich phenotype with a *H3.3* mutation. However, the mutation was present in both osteoclast-rich and osteoclast-deficient areas. Further research is required to explain the genotype–phenotype association between *H3F3A* p.G34W and osteoclast recruitment. This is only the second time that the p.G34W alteration has been associated with a malignant bone tumour. The first was reported in an osteosarcoma but no histological or clinical details of this case were reported [3]. We have previously reported 2 osteosarcomas harbouring a *H3.3* mutation, namely p.G34R in *H3F3A* and in *H3F3B*. Taken together these findings indicate that malignancy in bone tumours cannot be excluded on detection of *H3.3* mutations. In contrast, diagnosis of a GCT in the absence of a *H3F3A* alteration should be made with considerable caution in view of the high prevalence of *H3F3A* mutations in GCT, 95% (132/139 from current and previous study).

The role of *H3.3* genes in human neoplasms has been brought to the fore by large scale sequencing studies undertaken on paediatric brain tumours [18]. One of the most intriguing aspects of these mutations is the specificity of mutations for age and anatomical location of these tumours. Similarly, our work presented here confirms our previous observation of tumour type specificity of *H3.3* mutations in further bone tumour subtypes. Mutations at p.G34 were confined to tumours harbouring cells of the osteoclastic lineage, whereas mutations in p.K36 occurred solely in bone tumour of chondroblastic lineage. Furthermore, we found that mutations of *H3F3B* at G34 are exceedingly rare and have to date only been described in a single case of osteosarcoma. Finally, the association of the rare p.G34L mutation with tumours of small bones, caused by a double substitution, is noteworthy. The point mutations of *H3F3A* in brain tumours localise close to sites of post-translational modification of p.K27 and p.K36 resulting in methylation of H3K27 (transcriptional repression) and H3K36 (transcriptional elongation and splicing). It is, therefore, likely that these mutations, accompanied by subsequent post-translational modifications, exert their oncogenic effects through altered gene expression profiles in specific cells and in a particular developmental context. This proposition is supported by *in vitro* experimental evidence using glioma cell lines showing a clear link between histone mutation, methylation, gene expression and

tumourigenesis [18,19]. The specific roles that *H3.3* p.G34 mutations may have in tumourigenesis are still not known.

Kato Kaneko *et al* recently reported a high prevalence of *IDH2* alterations in GCT (16/20 cases): 13 were detected by direct Sanger sequencing and 3 by subcloning the PCR product and subsequent sequencing [9]. By contrast, we did not find any *IDH1* or *IDH2* mutations in the majority of 84 GCT and 22 osteoclast-rich tumours of the small bones of the hands and feet. A canonical *IDH2* mutation was detected in one of our cases but, on histological re-evaluation, we considered this case to represent a dedifferentiated chondrosarcoma. Although a rare finding, this could soon have therapeutic implications in view of the development of *IDH1/2* inhibitors, and a vaccine that targets mutant *IDH1* [20,21], and therefore, this case highlights the utility of genotyping tumours for diagnostic purposes. The reason for the findings by Kato Kaneko *et al* is unclear: we consider it is unlikely to be explained by a different cohort of tumours in view of the numbers of cases that we have analyzed. It is unlikely to be a sensitivity issue as the Ion Torrent platform is very sensitive. DNA contamination is a possibility, particularly if subcloning of the *IDH1/2* genes has been performed previously in this laboratory.

The presence of *H3F3A* p.G34 alterations in 96% of GCT along with their absence in other benign bone tumours including CB, osteoid osteoma, osteoblastoma, tenosynovial giant cell tumour and chondromyxoid fibroma provides evidence that *H3F3A* p.G34 mutations are a highly sensitive and specific marker for GCT. It is particularly valuable for distinguishing between those tumours with mutually exclusive alterations such as GCT and CB, and GCT and ABC, which harbour *USP6* fusion genes in 75% of cases [5,22]. In contrast, there is no genetic alteration characteristic of NOF. Although *H3F3A* p.G34W alterations were detected in 4/66 cases, we propose that these cases represent misdiagnoses, based on a multidisciplinary review of these cases. Indeed, in all cases studied here in which genotype was discordant with the original pathological diagnosis, we reviewed histology and imaging. In all but one case, we altered our original diagnosis in favour of the genetic findings. Nevertheless, using the detection of *H3.3* alterations in clinical practice would be more robust, if mutually exclusive markers were available for each of these tumour types, or if a second recurrent alteration was present in these tumours.

GCG of the small bone of the hands and feet and GCG of the jaws are considered to be distinct from conventional GCT (WHO classification [1]). It has been argued that these lesions represent a ‘solid

variant' of a primary ABC [23]. Agaram *et al* recently provided molecular evidence for this proposition by demonstrating *USP6* rearrangements in 8/9 GCG of the hands and feet [4]. They also reported no *USP6* alterations in eight conventional GCT, two of which were located in the finger. The combined data from our current and previous study [2] provide further evidence for the occurrence of solid variants of ABC in the bones of the hands and feet, and show for the first time that GCT also represents a significant number of osteoclast-rich tumours at these sites, highlighting the benefit of using these mutually exclusive genetic alterations for distinguishing these entities.

It is noteworthy that neither *H3.3* nor *USP6* alterations were detected in any of the 78 GCG of the jaw in our study indicating that GCG of the jaws are distinct from both GCT and solid variants of primary ABC. From a clinical perspective, it is important to note that osteoclast-rich lesions of the jaw also occur in patients with Noonan syndrome, neurofibromatosis (NF1) and cherubism (*SH2BP2*), and that these should be excluded before provision of a diagnosis of GCG [24,25]. Not all GCG included, in this study had been analysed for these alterations and it is, therefore, possible that some of the cases diagnosed as GCG represent syndrome-related lesions.

The introduction of screening of bone tumours for *H3.3* alterations along with other common bone cancer genes including *IDH1/2* alterations in cartilaginous tumours [8,9], *GRM1* rearrangements of chondromyxoid fibromas [26] and *USP6* alterations in ABC [5,6,22] adds to the armoury with which pathologists can provide accurate diagnoses. Although *H3.3* alterations does not stratify patients to therapeutic agents at this time, in the future this may be the case [20,21]. The study also highlights the changes required in laboratory practices if these new findings and those yet to be discovered are to be introduced effectively into clinical medicine.

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Author contributions

The concept was conceived by AMF. Review of the pathology was undertaken by histopathologists AMF,

MFA, RT, DB and GJ. Genome sequencing was performed at the CR-UK Oxford Molecular Pathology Institute (ABH) and analysed at Wellcome Trust Sanger by DW, PVL, SB, PT, and PC. Preparation of the samples and DNA was performed by BK. The experimental design was performed by NP, MFA, RH, AMF and DB. Analysis of the data was performed by NP, MFA, and AMF. The MS was written by AMF, DB, SB, NP and MFA.

Sequencing data have been deposited at the European Genome-Phenome Archive (EGA, <http://www.ebi.ac.uk/ega/>), which is hosted by the European Bioinformatics Institute (EBI); accession number pending.

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SUPPLEMENTARY MATERIAL ON THE INTERNET

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The following supplementary material may be found in the online version of this article:

Supplementary methods:

Table S1. Samples of giant cell tumour (GCT) and chondroblastoma (CB) with H3.3 alterations published in the previous study [2] with additional results to the original table published for USP6 analysis and targeted sequencing.

Table S2. Samples with diagnoses, clinical and genetic data from this study.

Table S3. Sample type and distribution.

Table S4. Primer sequences and description of the hot-spot targeted with PGM sequencer.

Table S5. Somatic substitutions and indels.

Figure S1. Genome-wide copy number profile of five giant cell tumours of bone. 4/5 tumours show essentially no copy number changes. Bottom: PD7524a harbours telomeric LOH in chromosomes 5q and 8p (arrows) assessed by Affymetrix SNP6. The smaller changes seen are likely to represent ‘noise’ as the sample was shown to be of low cellularity (33%). X-axis: genomic position (ascending). Y-axis: copy number. Numbers in plot refer to chromosome. Red and green: major and minor allele, respectively.

Figure S2. (A) Photomicrographs of a tumour presenting in a 59 year old originally diagnosed as a GCT. The biopsy showed a fragment of mature hyaline cartilage and an osteoclast-rich mitotically active tumour. The tumour was wild type for H3F3 and an *IDH2* alteration was detected. Following re-evaluation and in the context of the genetics a diagnosis of a dedifferentiated chondrosarcoma was made. (B) Radiograph and photomicrographs of subarticular lytic tumour in the proximal tibia. The imaging suggested a GCT but the histology showed a mitotically active spindle cell tumour with epithelioid differentiation, supported by the cytokeratin expression (inset) and an osteoclast-rich area. The tumour was classified as an unusual high grade osteoclast-rich tumour with adamantinoma-like features.

Figure S3. Algorithm-based schema illustrating the approach to reaching a diagnosis of osteoclast-rich tumours.