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Discovery of human-similar gene fusions in canine cancers

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Abstract

Canine cancers represent a tremendous natural resource due to their incidence and striking similarities to human cancers, sharing similar clinical and pathological features as well as oncogenic events including identical somatic mutations. Considering the importance of gene fusions as driver alterations, we explored their relevance in canine cancers. We focused on three distinct human-comparable canine cancers representing different tissues and embryonic origins. Through RNA-Seq, we discovered similar gene fusions as those found in their human counterparts: $IGK-CCND3$ in B-cell lymphoma, $MPB-BRAF$ in glioma, and $COL3A1-PDGFB$ in dermatofibrosarcoma protuberans-like. We showed not only similar partner genes but also identical breakpoints leading to oncogene overexpression. This study demonstrates similar gene fusion partners and mechanisms in human-dog corresponding tumors and allows for selection of targeted therapies in preclinical and clinical trials with pet dogs prior to human trials, within the framework of personalized medicine.
Introduction

During the last decade, pet dogs have arisen as a relevant but under-used model for cancer genetics and therapies (1). Although extremely useful for cancer research, rodent models show limitations in their ability to fully reproduce the complexity of spontaneously occurring human tumors (2). Since canine tumors address part of these concerns, there is a growing interest in using this “natural cancer resource” in translational cancer research for both human and dog medicine benefit. Inclusion of pet dogs in preclinical and clinical trials should accelerate the screening of new treatments by reducing the time and cost of evaluating efficacy, pharmacokinetics and toxicity (3). These advantages have already justified ongoing trials with pet dogs (4).

To use canine cancers as models for the development of new treatments, it is essential to determine human and canine tumors correspondences. The physiopathology of canine tumors presents strong similarities with human tumors in their biological behavior, histopathological features and response to treatments (2). Moreover, comparative oncology studies show that some canine and human cancers share similar genetic aberrations including chromosomal instability, common cytogenetic aberrations, involvement of identical oncogenes or tumor suppressors, and even identical somatic mutations (5). The identification of such driver genetic alterations, both in dogs and humans, are highly informative for the development of dedicated new targeted therapies such as the development of c-kit inhibitors for canine mast cell disorders (6). Among such driver events, we focused on translocations that generally lead to overexpression of oncogenes. In humans, these recurrent translocations have been described in hematological disorders, sarcomas and carcinomas; most of them are
specific, and even pathognomonic for tumor types and represent precious tools for diagnosis and relevant therapeutic targets (7,8). Although well described in humans, such translocations in dogs have only been explored and identified by fluorescence *in situ* hybridization (FISH) to date, in two canine hematological tumors: a chronic lymphocytic leukemia with a \( \text{BCR-ABL} \) fusion and a Burkitt’s lymphoma with a \( \text{IGH-MYC} \) translocation (9).

In the present study, we aim to demonstrate that pet dogs are relevant spontaneous models for human oncology with conserved key gene fusions, not restricted to hematological disorders by identifying chimeric transcripts with new and highly resolutive methods (NGS). Thus, we used whole transcriptome sequencing (RNA-Seq) of cancers with different origins to allow the detection of gene fusions without preconceived candidates. We report here, for the first time, chimeric transcripts in canine cancers reflecting chromosomal translocations that are comparable to human translocations in: a dermatofibrosarcoma protuberans-like (DFSP-like); an oligodendroglioma; and a diffuse large B-cell lymphoma (DLBCL). We then validated these gene fusions on DNA and RNA from the tumors, providing the breakpoint positions at the single-nucleotide resolution. We finally showed that similarly to the orthologous human cancers, these canine cancers not only share identical breakpoints but also overexpression of the targeted oncogenes. These striking molecular similarities between human and canine cancers pave the way for the use of the canine model in targeted therapies for the benefit of both human and dog patients.

**Material & Method**

*Samples*
Blood and tissue biopsy samples from dogs were collected by a network of veterinarians through the Cani-DNA biobank (http://dog-genetics.genouest.org), which is part of the CRB-Anim infrastructure. The work with dog samples was approved by the CNRS ethical board, France (35-238-13) for UMR6290 and dog owners consented to the use of data for research purposes anonymously. We selected 3 distinct cancers (lymphomas, sarcomas and gliomas) representing different tissues (neurological, subcutis, hematological) and different embryonic origins, with known fusions in the human counterpart. The set of samples included 3 lymphomas (2 DLBCLs and 1 T lymphoma), 2 gliomas and 1 dermatofibrosarcoma protuberans-like.

Histological diagnosis

The diagnosis of the six tumors was re-evaluated by dedicated veterinary (JA) and human pathologists (JMC and AR for the DFSP-like and the glioma respectively). Immunohistochemical staining was performed using antibodies showing reactivity with the dog’s protein according to the manufacturer or to biocompare website: anti CD34 (Santa Cruz Biotechnology Dallas, TX, Ref SC-7045) and anti PDGFB (Novus Biological, Littleton, CO, Ref NBP1-52533), anti BRAF (Abbiotec, San Diego, CA, Ref 251460), anti CCND3 (Aviva, San Diego, CA, Ref AVARP03038_P050) for the DFSP-like, the glioma and DLBCL respectively. Appropriate negative controls were performed with normal immune serum of rabbit for CCND3 and BRAF, of goat for PDGFRB.

DNA and RNA isolation

The germline DNA from blood, DNA and RNA from tissues were isolated using respectively the Blood L, Tissue and RNA II NucleoSpin® kits (Macherey-Nagel, Duren, Germany) according to the manufacturer's instructions.
**RNA-Seq sample preparation and sequencing**

RNA-Seq was performed using the GIGA genomics facility. Briefly, Illumina Truseq RNA Sample Preparation kit V2 was used to prepare libraries from 1µg of total RNA with a RNA Integrity Number greater than 7.3 (Agilent Technologies, Santa Clara, CA). Poly-A RNAs were purified with polyT-coated magnetic beads and chemically fragmented to a median size of 200 nucleotides. These were used as a template for first strand synthesis in the presence of random hexamers and second strand synthesis afterwards. Next, double stranded cDNA ends were end-blunted and adenylated at 3’OH extremities before ligation to adaptors containing the indexes. Finally, the adapter-ligated library fragments were enriched by PCR following Illumina’s protocol and purified with Ampure XP magnetic beads (Agencourt, Beverly, MA). Final libraries were validated on a Bioanalyser DNA 1000 chip (Agilent Technologies, Santa Clara, CA) and quantified by qPCR with the KAPA library quantification kit (Kapa Biosystems, Wilmington, MA). Sequencing was performed on an Illumina HiSeq 2000 instrument using the PE 2X100 cycles protocol.

**Identification of fusion transcripts**

Candidate gene fusion transcripts were identified with CRAC and chimCT softwares ([http://cractools.gforge.inria.fr/](http://cractools.gforge.inria.fr/)) using the CanFam3 canine genome reference. Fusion transcripts were manually filtered by discarding transcripts with less than five reads spanning the fusion junction and three reads encompassing the two fusion transcripts, as well as transcripts involving paralogous genes or pseudogenes, transcripts identified as read-through, transcripts with no Ensembl annotation (Supplementary Table 1).
RT-PCR

Reverse transcription was performed on 1 μg of total RNA from tumor or healthy tissue using the High-capacity cDNA Reverse Transcription kit (Applied Biosystems, Foster City, CA) according to the manufacturer's instructions.

Validation of the fused transcript

PCR amplifications were performed on cDNA samples diluted 1:40 using the Type-it Microsatellite PCR kit (Qiagen, Hilden, Germany) with dedicated primers (Supplementary Table 2). PCR products were sequenced with a 370 ABI sequencer using the BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems, Foster City, CA). The presence of fusion transcripts was assessed by alignment of sequences on the CanFam 3 reference dog sequence (https://genome.ucsc.edu/).

Breakpoint mapping on genomic DNA

PCR amplifications were performed on DNA to validate fusion transcripts. Primers specific for each breakpoint (Supplementary Table 2) were designed on CanFam3 to span the candidate introns flanking the identified exons in the chimeric transcripts. All combinations between forward and reverse primers from translocated chromosomes were tested. The large regions of about 1 kb were amplified from 10 ng/µl diluted DNA samples using the Phusion Hot Start II DNA Polymerase kit (Thermo scientific, Waltham, MA).

Exon expression analysis RT-PCR

qRT-PCR was performed on cDNA samples diluted 1:20 with the SYBR Green PCR master mix on the 7900HT Fast Real-Time PCR System (Applied Biosystems, Foster City, CA).
CA) using standard procedures. Each PCR was carried out in triplicate. Relative amounts of the transcript were determined using the delta-delta Ct method. Canine HPRT gene (ENSCAFG00000018870) for the lymphoma and the DFSP-like, and TBP gene (ENSCAFG00000004119) for the glioma, were used as housekeeping genes. The mRNA levels in each tumor were calculated as a fold increase compared with at least 6 different healthy tissue mRNA controls: 6 healthy skin samples, 8 healthy cortex samples, 6 healthy lymph nodes for the fibrosarcoma, glioma and lymphoma respectively.

**Western Blot**

Cellular protein extracts were prepared using a cell lysis buffer containing 20mM Tris-HCl pH 7.5, 150mM NaCl, 1mM EDTA, 1mM EGTA and supplemented with 1mM PMSF, 1X EDTA-free cocktail protease inhibitor (Roche Diagnostic, Meylan, France), 30mM sodium fluoride, 40mM glycero phosphate, 1mM sodium orthovanadate, 0,5% Triton X-100. Protein concentrations were determined by the BCA protein assay (Sigma-Aldrich, St Louis, MO) using bovine serum albumin as a standard.

Protein samples were denatured for 10min at 95°C, and equal amounts of cell proteins (20 μg) were subjected to 10% SDS-PAGE and transferred onto nitrocellulose membranes (Amersham - GEH life, Arlington Heights, IL). Membranes were probed with suitable antibodies. The primary antibodies were: anti-BRAF (Ref 251460, 1:5000, Abbiotec, San Diego, CA) and anti-β-actin (BA3R, 1:5000, Millipore, Temecula, CA). Horseradish-Peroxidase-conjugated secondary antibodies were purchased from Jackson ImmunoResearch (Suffolk, UK). Signals were detected using the LAS-4000 Imager (Fuji Photo Film, Tokyo, Japan). siRNAs were purchased from IDT DNA (Coralville, IO):

- siBRAF1: 5’- CAUGAAGACCUCACAGUAAAAAUAG-3’,
- siBRAF2 5’- GACCAAAUUUGAGAUGAUAAAGCTT-3’,
siCTR (NC1) 5’-CGUUAUCGCGUAUAUACGCGUAT-3’

Two siRNAs targeting BRAF were designed to target two distinct regions of BRAF (nucleotides 1717-1740, 1599-1623 respectively for ENSCAFT00000006305) and ordered to IDT DNA. siRNAs were transfected into human melanoma cells 501Mel or into a canine oral melanoma cell line, kindly provided by Dr. D. Tierney, using Lipofectamine RNAiMAX (Life Technologies, Carlsbad, CA) following the manufacturer’s instructions. Three days later, BRAF expression levels were analyzed by western blotting experiments.

**FISH experiments**

MBP-BRAF fusion tests were performed on 4 µm sections of formalin-fixed paraffin-embedded tissue blocks using the bacterial artificial chromosome (BAC) clones 42G22-97M23 and 160H02-360E1 (http://bacpacresources.org/library.php?id=253). These BAC clones were labeled with green-dUTP (Abbott Molecular, Downers Grove, IL) and Cy3-dCTP (Amersham Biosciences, Buckinghamshire, UK) and hybridized proximal to MBP and distal to the BRAF breakpoint regions, respectively. Slides were analyzed by an experienced cytogeneticist (FC), using a fluorescence microscope (Axioskop2, Axios Imager Z2, Zeiss, Göttingen, Germany) and Isis imaging software (Metasystems, Altlussheim, Germany). Per case, at least 100 non-overlapping tumor nuclei were examined.

**Statistics**

All statistical tests were performed with the R software.

**Results and discussion**
Previous cytogenetic studies have underlined the importance of gene fusions in hematological disorders and sarcoma in humans (8). To investigate gene fusions in canine cancers, we analyzed 6 tumors belonging to 3 distinct cancers, representing different tissues (neurological, subcutis, hematological) and embryonic origins, and for which the human corresponding tumors are known to involve key gene fusions. We performed RNA-Seq, using dedicated softwares followed by classical filters and manual curation to exclude false-positive chimeras, we finally identified 3 gene fusions in 3 of the 6 tumors (Table 1 and Supplementary Table 1).

1-A COL3A1-PDGFB fusion in a canine dermatofibrosarcoma protuberans-like

In human dermatofibrosarcoma protuberans (DFSP), the translocation t(17;22) leading to the fusion of the Collagen type 1 alpha 1 (COL1A1) and Platelet-derived growth factor B chain (PDGFB) genes is considered as pathognomonic of DFSP (10). Thus, we analyzed a canine DFSP-like case (Supplementary Figure 1) and we found a chimeric transcript resulting from the fusion between the Collagen type 3 alpha 1 (COL3A1) and the PDGFB genes. Since COL3A1 and COL1A1 are paralogous genes, we anticipated that this canine translocation is similar to the human one. Indeed, as in humans, the breakpoint in PDGFB is localized in intron 1 (Figure 1a), moreover the fusion validation on the cDNA and the genomic DNA of this tumor (Figure 1b-d) shows a conserved reading frame between the first COL3A1 exons and the exon 2 of PDGFB. Similar to the situation in human DFSP (11), we hypothesized that this translocation, placing PDGFB under the regulation of the COL3A1 promoter, induces the overexpression of PDGFB in the canine DFSP-like tumor. This is confirmed by the expression levels of exons retained in the
chimeric transcript, which are increased on average 9.6-fold in comparison to healthy tissues (p-value <0.001, one-tailed Wilcoxon test) (Figure 1e). Moreover, PDGFB protein is detectable in the tumor by immunostaining (Supplementary Figure 1). Complementary to this, the alignment of canine and human PDGFB protein sequences (Supplementary Figure 2) shows the conservation of the 2 proteolytic cleavage sites in the canine PDGFB and in the putative chimeric protein resulting from \textit{COL3A1/PDGFB} fusion. Under these conditions, the canine \textit{COL3A1/PDGFB} should produce a chimeric precursor protein, transformed into a mature growth factor after proteolysis and forming an autocrine loop, as for the human \textit{COL1A1/PDGFB} fusion (11). Since the chimeric protein \textit{COL1A1/PDGFB} is targeted by imatinib for human DFSP (7,10), regarding the genetic and biological similarities between human DFSP and canine DFSP-like, we predict similarities in treatment responses. Finally, we propose the \textit{COL3A1} gene as a new candidate partner gene for human DFSP, like the recently discovered \textit{COL1A2} (12).

2-A \textit{MBP-BRAF} fusion in a canine glioma

Human gliomas are a heterogeneous group of tumors of the central nervous system with \textit{BRAF} alterations (somatic mutations or fusions) described almost in all subgroups (13). Therefore we analyzed two canine grade 3 anaplastic oligodendrogliomas (WHO classification) (Supplementary Figure 3). In one of them, we identified a relevant chimeric transcript resulting from the fusion between the \textit{Myelin Basic Protein (MBP)} and the \textit{B-Raf proto-oncogene (BRAF)} genes. Like in human gliomas with breakpoints localized between \textit{BRAF} intron 8 and 10 (14), the breakpoint of \textit{BRAF} in this canine glioma is within intron 7, orthologous to human \textit{BRAF} intron 8 (Figure 2a). This fusion
was validated on the cDNA and the genomic DNA of this tumor (Figure 2b-d). The reading frame is conserved between the first MBP exons and the last exons of BRAF. Since the MBP promoter is strongly active in glioma cells (15), we anticipated that this translocation could also induce BRAF overexpression. We confirmed this hypothesis by measuring the expression levels of exons retained in the chimeric transcript; there was an average 12.8-fold increase in comparison to healthy tissues (p-value = 0.0039, one-tailed Wilcoxon test) (Figure 2E). Moreover, BRAF protein is detectable in the tumor by immunostaining and a fused protein is detectable in the tumor sample by western blot (Supplementary Figure 3). Since the alignments of the canine and human BRAF protein sequences showed that the canine chimeric protein has lost its N-terminal auto-inhibitory domain (Supplementary Figure 4), this alteration should induce a constitutive BRAF kinase activity as shown in human cancers (16). In human gliomas, BRAF is mainly fused with KIAA1549 but other partners have been described (14), while most of these fusions are found in astrocytomas, they are also reported in oligodendrogliomas (17). We propose MBP as a new candidate partner for BRAF fusions in human gliomas. For these human tumors, with limited therapeutic options, BRAF alterations offer new therapeutic strategies and clinical trials are ongoing (16). In this context, spontaneous canine glioma with BRAF fusions provides a unique model to develop new and more efficient treatments.

3-An IGK-CCND3 fusion in a canine B-cell lymphoma

In human lymphomas, alterations during the natural recombination process of immunoglobulin genes lead commonly to fusions with oncogenes associated with their overexpression. Thus we analyzed 3 canine lymphomas (two DLBCLs, and one T-cell
lymphoma) and found one aberrant transcript revealing a fusion between the *Immunoglobulin light chain kappa locus* (*IGK*) with *Cyclin-D3* (*CCND3*) in one DLBCL sample (Table 1) (Figure 3a). This fusion was validated on the cDNA and the genomic DNA of this tumor (Figure 3b-d). Such translocations involving *CCND3* and immunoglobulin genes have already been described in human B-cell malignancies including DLBCL, which could explain the overexpression of *CCND3* found in 10% of human DLBCL (18). Thus, we anticipated that the canine translocation also induces the overexpression of *CCND3*. This is supported by the fact that *CCND3* is overexpressed in canine DLBCL (Figure 3e) and the CCND3 protein is easily detectable by immunohistochemistry in the canine DLBCL (Supplementary Figure 5). While, *IGK-CCND3* fusion was not identified in human lymphomas, overexpression of *CCND3* driven by *IG*-mediated translocations are expected and searched in cases showing break-apart probes for *CCND3*. Moreover fusions involving *IGK* and other cyclins, of which *CCND2*, have already been detected (19). In addition, the overexpression of *CCND3* in human DLBCL is a prognostic factor associated with poor clinical outcome (20). Thus, canine lymphomas with fusions involving cyclin D could be used for clinical trials targeting cyclins.

**Conclusion**

Our work reveals similar fusions in corresponding cancers between dogs and humans and identifies for the first time chimeric transcripts involving the same fused oncogenes with similar rearrangements leading to malignancy. The identification of these three fused transcripts in three different tumors supports the existence of other fused transcripts in other subtypes of canine cancers, similar to what is found in humans. With
NGS methods, which are revolutionizing the identification of gene fusions without preconceived idea of partner genes, such canine models will benefit the scientific community, allowing development of new targeted therapies. In the context of the “One Health” concept, such results of comparative oncology lead us to anticipate that human and veterinary medicine will benefit from clinical trials that include pet dogs as patients.
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References

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**Table 1:** Fusions identified by RNA-Seq in the canine tumors. Partner genes and chromosomal positions are indicated for the three tumors with the number of reads spanning the fusion junction and encompassing the two fusion transcripts. CFA: *Canis familiaris* Autosome.
Figures Legends

**Figure 1:** Characterization of the COL3A1-PDGFB fusion. (A) Scheme of the t(10;36). Exons of COL3A1 (ENSCAFT00000047312) and PDGFB (ENSCAFT0000002101) are depicted as well as the breakpoint positions on CFA36 (blue) and CFA10 (red), indicated with . CFA: *Canis familiaris* Autosome. (B) Sequence of the breakpoint on genomic DNA. The 3 nucleotides flanking the dotted line map to both chromosomes. (C) Electrophoresis of PCR products detects the translocation in tumor but not in the germinal blood DNA of this case. (D) Scheme of the transcription resulting from the t(10;36): the sequence of the tumor cDNA reveals the expression of one fusion transcript involving exon 27 of COL3A1 and exon 2 of PDGFB. (E) Relative expression levels of PDGFB exons in the tumor in comparison with 6 healthy skin samples (* p-value < 0.05 one-tailed paired-samples Wilcoxon test). Expression levels of exons 2 to 6 show overexpression of PDGFB exons retained in the chimeric transcript. The PDGFB wild type transcript expression, represented by expression levels of exons 1-3, shows no significant difference in the tumor in comparison with an expected value of 100% (p-value = 0.31, two-tailed one sample Wilcoxon test). These results indicate that the overexpression of PDGFB exons in the chimeric transcript is the consequence of the fusion.

**Figure 2:** Characterization of the MBP-BRAF fusion. (A) Scheme of the t(1;16) translocation. Exons of MBP (ENSCAFT000000024) and BRAF (ENSCAFT0000006305) are depicted as well the breakpoint positions on CFA1 (blue) and CFA16 (red) indicated with . CFA: *Canis familiaris* Autosome. (B) Sequence of the breakpoint on genomic DNA. The 3 nucleotides flanking the dotted line map to both chromosomes. (C) Electrophoresis of the PCR product detecting the translocation in tumor DNA but not on germinal blood DNA of this case. (D) Scheme of the transcript resulting from the t(1;16) translocation: the sequence of the tumor cDNA reveals the expression of one fusion transcript involving the exon 1 of MBP with exon 8 of BRAF. (E) The histogram shows the relative expression levels of BRAF exons in the tumor in comparison with 8 healthy brain samples. The fusion induced overexpression of the retained exons 11-13 on average 12.8-fold (p-value ≤ 0.01, one-tailed Wilcoxon test) in comparison with healthy canine brain tissues. The retained exons are overexpressed in
comparison to exons localized in 5’ of the breakpoint (** p-value ≤ 0.01, two-tailed paired two-Sample t-test). Expression levels of exons 7-8, flanking the breakpoint, showed that the BRAF wild type transcript is significantly under-expressed in the tumor (p-value ≤ 0.01, one-tailed Wilcoxon test). These results showed that the overexpression of BRAF exons retained in the chimeric transcript is linked to the fusion.

**Figure 3:** Characterization of the IGK-CCND3 translocation in a canine DLBCL. (A) Schematic representation of the t(12;17) translocation. Exons of CCND3 (ENSCAFT00000002538) and IGK (ENSCAFT00000011790) are depicted as well as the breakpoint positions on CFA12 (red) and CFA17 (blue), indicated with . CFA: Canis familiaris Autosome. (B) Sequence of the breakpoint on genomic DNA. (C) Electrophoresis of the PCR product detecting the translocation in the tumor DNA but not in the germinal blood DNA of this case. (D) Scheme of the transcription resulting from t(12;17) translocation: the sequence of the tumor cDNA reveals the expression of one fusion transcript involving a not yet annotated exon of CFA12 with exon 5 of IGK. (E) RT-qPCR experiments show the overexpression of the CCND3 mRNA transcript in the tumor, on average 13-fold (p-value < 0.05 one-tailed One-Sample Wilcoxon test) and 22-fold (**** p-value < 10^{-4} one-tailed Two-Samples Wilcoxon test), as compared with 6 healthy lymph nodes and 6 other DLBCLs respectively.