

Debunking the Myth of the Endogenous Antiangiogenic Vegfaxxxb Transcripts

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1	Debunking the myth of the endogenous anti-angiogenic Vegfaxxxb transcripts.
2	
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16	DS edited the MS and contributed additional data analysis; all authors approved the final draft.
17	
18	Abstract
19	In this article we critically assess evidence for the existence of a family of anti-angiogenic
20	Vegfaxxxb transcripts, arising from the use of a phylogenetically conserved alternative distal
21	splice site within exon 8 of the VEGFA gene. We explain that prior evidence for Vegfaxxxb
22	transcripts in tissues rests heavily upon flawed RT-PCR methodologies, with the extensive use of
23	5'-tailing in primer design being the main issue. Furthermore, our analysis of large RNA-seq
24	datasets (human and ovine) fails to identify a single Vegfaxxxb transcript. Therefore, we

challenge the very existence of *Vegfaxxxb* transcripts, which further questions the physiological

relevance of studies based on the use of "anti-VEGFAxxxb" antibodies. Our analysis has
implications for the proposed therapeutic use of isoform-specific anti-VEGFA strategies for
treating cancer and retinopathies.

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30 Classical angiogenic VEGFA isoforms

VEGFA is a key regulator of vascular homeostasis and therapeutic target in pathological 31 angiogenesis. The VEGFA gene is subject to alternative splicing, which occurs primarily in 32 exons 6 and 7 (Figure 1) [1-4]. In humans, the most abundant mature translated isoform is 33 VEGFA165 (i.e. this isoform comprises 165 amino acids), which lacks exon 6. Three other 34 35 isoforms are present in moderate/high amounts: VEGFA121, which lacks both exons 6 and 7, VEGFA189, which includes exon 7 but uses an alternative 5'-donor site in exon 6 and 36 VEGFA206, which includes both exons 6 and 7 (Figure 1B & Figure 1E). These VEGFA 37 38 isoforms exist in other species, including mouse, sheep and cattle, but they are one amino acid shorter than their human counterparts. Other less abundant isoforms have been described, some 39 of which are also generated through alternative splicing within exons 6/7 (VEGFA145, 40 VEGFA148, VEGFA183). The amino acid stretches encoded by exons 6 and 7 are enriched in 41 clusters of basic amino acids, which confer VEGFA with the ability to bind the extracellular 42 matrix and thus mitigate its diffusion. Consequently, VEGFA121 is highly diffusible, while 43 VEGFA189 and VEGFA206 remain bound to the extracellular matrix and VEGFA165 has 44 intermediate matrix binding characteristics [1-4]. VEGFA isoforms bind with varying affinities 45 to the tyrosine kinase receptors VEGFR1 and VEGFR2 and recruit the co-receptor Neuropilin 1 46 to activate intracellular signaling pathways as illustrated on the left side of Figure 1G [5,6]. 47 Critically, all VEGFA isoforms promote angiogenesis, which is key to the progression of 48 tumorigenesis [7]. 49

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53 Non-classical anti-angiogenic VEGFAxxxb and VEGF-Ax isoforms

In 2002, Bates and colleagues [8] reported a novel human Vegfa splice variant, generated 54 through use of an alternative distal splice site within the last exon of the VEGFA gene, exon 8 55 (see Figure 1C and Figure 1D). The transcript was dubbed *Vegfa165b*. Follow-up studies 56 suggested that the encoded VEGFA165b protein product belongs to an entire VEGFAxxxb 57 family of proteins, which are derived from transcripts that use the distal splice site in exon 8 58 (Figure 1E) and bear anti-angiogenic properties [3,9-14]. From a functional standpoint, 59 60 VEGFA165b was shown to bind to VEGFR2 with the same affinity as VEGFA165 [15-18] but failed to bind to Neuropilin 1 as illustrated on the right part of Figure 1G [16,17,19]. 61 VEGFA165b induces VEGFR2 tyrosine phosphorylation in a dose-dependent manner, albeit less 62 efficiently and more transiently than VEGFA165 [15-17,19]. However, this impaired ability of 63 VEGFA165b to activate intracellular signaling pathways may not apply to all cell types [20,21]. 64 Another isoform, dubbed VEGF-Ax, has recently been described [22]. VEGF-Ax arise from 65 programmed translational read-through, such that the stop-codon of classical VEGFA isoforms 66 leads to insertion of a Serine residue, and the in-frame stop-codon of putative VEGFAxxxb 67 isoforms is used instead. In other words, VEGF-Ax and VEGFAxxxb isoforms would share the 68 same C-term sequence. However, the in vivo existence of VEGF-Ax is disputed and in vitro data 69 demonstrate pro-angiogenic properties [23]. 70

71

72 Existence of VEGFAxxxb isoforms: clinical implications

Overall, VEGFA165b appears to behave as a partial agonist [16], which competes with
VEGFA165 for VEGFR2/Neuropilin 1 binding. This may explain the reduced angiogenic
potential of VEGFA165b [20,23,24]. Based on this, Harper & Bates proposed a model in which

the balance between endogenous levels of angiogenic VEGFAxxx and anti-angiogenic 76 77 VEGFAxxxb isoforms sets the angiogenic potential of a tissue [3]. They further proposed that VEGFAxxxb isoforms predominate in many normal cells and tissues where they can amount to 78 more than 50% of total VEGFA protein [10]. In normal colonic tissues in particular, 79 VEGFAxxxb variants have been reported to comprise as much as 95% of all VEGFA [25]. The 80 proposed use of the alternative distal splice site within exon 8 is therefore not a rare event, and 81 its prevalence implies major physiological relevance. However, in pathological conditions such 82 as cancer and retinopathies, VEGFAxxxb levels appear to be substantially downregulated. 83 Considering the crucial role of VEGFA in angiogenesis of various cancers and retinopathies 84 85 [1,26-28] such a finding may have broad clinical implications.

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Indeed, current strategies to target VEGFA, such as the anti-VEGFA antibody bevacizumab, 87 88 Avastin®) or aflibercept (VEGFA-trap), would indiscriminately inhibit both angiogenic VEGFAxxx and anti-angiogenic VEGFAxxxb isoforms. Bates et al showed that targeting 89 90 VEGFA in colorectal cancer was more efficient if endogenous levels of VEGFAxxxb were low and proposed that assessment of the ratio of VEGFAxxxb to VEGFAxxx could potentially 91 predict response to bevacizumab and other therapies directed against VEGFA [3]. Therefore, 92 modification of VEGFA splicing to alter the VEGFAxxxb/VEGFAxxx balance, or development 93 of compounds to target only VEGAxxx isoforms, may have therapeutic value for treating 94 cancers and retinopathies [29]. In 2009, D.O. Bates and S.J. Harper filed a patent for the "Novel 95 use of VEGFAxxxb", which covers the use of these isoforms (primarily VEGFA165b) for 96 several potential therapeutic applications. 97

98

99 The Controversy: Do VEGFAxxxb isoforms really exist?

Despite a substantial amount of literature reporting evidence for the existence of VEGFAxxxb 100 101 isoforms, the VEGFA scientific community is divided on this issue. We and others have yet to detect these isoforms in normal and pathological tissues and have thus concluded that 102 103 VEGFAxxxb isoforms, if they do exist, are likely not of physiological relevance. In recent reviews on VEGFA signaling and disease, VEGFAxxxb biology is largely disregarded [7] or 104 105 notably overlooked [30]. Until now, evidence for the existence of VEGFAxxxb isoforms has 106 been provided through PCR-based approaches and studies using an antibody generated against the sequence TCRSLTRKD encoded by putative exon 8b of the human Vegfa gene. Here, we 107 critically review and reassess the evidence which led to the assumption that "there is an 108 important role for VEGFAxxxb isoforms in normal physiology" [10]. Our own findings [31-33] 109 and a thorough analysis of the literature has led us to question and re-evaluate the existence of 110 *Vegfaxxxb* transcripts 111

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113 Absence of Evidence: Detection of *Vegfaxxxb* transcripts with flanking primers.

Since alternative splicing affects exons 6, 7 and 8, the best way to identify all Vegfa splice 114 variants is to use primers flanking these regions. Such a primer design is expected to amplify all 115 Vegfa isoforms. To do so, the forward primer has to be located within exons 1-5 (common to all 116 splice variants) and the reverse primer located downstream of the putative exon 8b splice site. If 117 Vegfaxxx and Vegfaxxxb splice variants are sister families and Vegfaxxxb mRNAs represent a 118 large fraction of total Vegfa mRNA, at least in normal tissues [10], a standard RT-PCR with the 119 aforementioned primers would be expected to yield at least 6 main products/bands corresponding 120 to mRNA for the major isoforms VEGFA189, VEGFA165 and VEGFA121 and their "sister 121 bands" for VEGFA189b, VEGFA165b and VEGFA121b, albeit at different abundances. The 122 bands encoding *Vegfaxxb* isoforms would be 66bp shorter than their respective counterparts 123 (see Figure 1D) thereby yielding 3 "doublets" easily discernible on an agarose gel. However, in 124

their seminal paper reporting on the discovery of Vegfa165b [8], Bates et al stated that "PCR of 125 the full-length product using primers V165K (complementary to the translation initiation site of 126 the other isoforms of VEGF) and V165X (a primer downstream of the original 3'-UTR) resulted 127 in one strong band at ~670bp". Considering the strategy used, it is difficult to explain why other 128 Vegfaxxx and Vegfaxxxb transcripts were not identified by this initial study. Indeed, as mentioned 129 above, at least 6 bands, corresponding to the 3 "doublets" (VEGFA189/189b, VEGFA165/165b 130 and VEGFA121/121b), should have been observed by agarose electrophoresis of PCR products. 131 Furthermore, nested PCR with a 3'-UTR and exon 7a primers resulted in a strong band at 132 ~130bp confirming that the full-length was VEGFA165b". 133

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We and others have adopted this strategy in order to identify Vegfaxxxb transcripts in human, 135 mouse, sheep and cattle but failed to identify Vegfa165b mRNA - or other Vegfaxxxb products 136 [22,31,33,34]. Instead, only three bands were observed (see Figure 1B and Figure 1F) and 137 sequencing of the PCR products showed unequivocally that they encoded VEGFA189, 138 139 VEGFA165 and VEGFA121 [22,24,32-34]. Therefore, an unbiased and straightforward RT-PCR strategy to simultaneously amplify all Vegfa isoforms failed to support the existence of 140 Vegfaxxxb mRNA. Since these flanking primers could not discriminate Vegfaxxx from putative 141 Vegfaxxxb transcripts, isoform-specific primers were developed. 142

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144 Detection of *Vegfaxxxb*-like transcripts with "isoform-specific primers": a tale of a tail.

Over time, there has been a considerable drift in the design of primers aimed at the specific identification of *Vegfa165b* that span the exon 7 – exon 8b splice site. However, in most studies, the exon 7-specific stretch has been greatly lengthened at the expense of the exon 8b-specific region (see **Figure 2A** and **Table S1**). Since exon 7 would be shared by both *Vegfa165* and *Vegfa165b*, the isoform-specificity of these reverse primers is questionable. Technically, the

necessary and sufficient primer design to allow specific detection of Vegfa165b mRNA would 150 include the shortest possible 3'-anchor within exon 7; based on standard PCR principles (see 151 Figure 3) one base specific to exon 7 on the 3'-end of the primer would suffice. However, in the 152 case of the human sequence, since the AT motif would be shared by both Vegfa165 and 153 Vegfa165b (see Figure 1) this means that the last 3bp at the 3'-end of such a primer would 154 anneal to exon 7 with the last nucleotide discriminating between Vegfa165 and Vegfa165b 155 156 isoforms (see Figure 2A, Primer P1). Surprisingly, this most simple strategy has never been used in the literature. 157

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159 Rather than using such a "minimal and sufficient" primer design for detection of Vegfaxxxb isoforms, the design of the primer has been "adapted". The initial design, published in the 160 seminal Vegfa165b paper by Bates et al [8], comprised of 7bp complementary to either Vegfa165 161 162 or Vegfa165b (Figure 2A, Primer P2). This primer has been used in a number of subsequent studies by the same group [15,35-39] and other research teams that used RT-PCR to demonstrate 163 the existence of Vegfaxxb transcripts [40,41]. In 2007, Ribeiro et al used a modified version of 164 this primer for their studies in pig [42], with 9bp complementary to either Vegfa165 or 165 Vegfa165b (Figure 2A, Primer P3), while Baba et al used 8bp complementary to either 166 Vegfa165 or Vegfa165b [43]. Overall, the number of bases complementary to either Vegfa165 or 167 Vegfa165b has been on the rise since 2007: 13 bp in 2008 ([44]; Figure 2A, Primer P4), 14bp in 168 2009 (study in rat; [45]) and 15bp in 2010-2011 ([46,47]; Figure 2A, Primer P5). Finally, a 169 design including no less than 17bp common to both Vegfa165 or Vegfa165b was proposed to 170 specifically amplify *Vegfa165b* mRNA ([48]; Figure 2A, Primer P6). We also note that the same 171 methodology has been used for the design of a Vegfa165b-specific forward primer, which 172 includes 14bp that would also anneal to *Vegfa165* ([34]; Figure 2A, Primer P7). This approach 173 has also been used by others to "specifically" detect *Vegfa121b* isoforms ([29,44]; see Table S1). 174

We demonstrated that this primer design is responsible for PCR artefacts, due to 5'-tailing, that 176 could be mis-interpreted as *Vegfaxxxb* transcripts ([31,33]; see Figure 3). The conclusion of 177 these studies was that PCR products correspond to putative Vegfaxxxb transcripts only because 178 primers were designed to include on their 5'-end a few bases of the Vegfaxxxb sequence itself. 179 This method has been used for decades, most often to add restriction sites to aid cloning. Good 180 examples are provided by Catena et al [20] and Ganta et al [49], who flanked the 5'-end of their 181 reverse primers with an *HindIII* (or *BamHI*, respectively) restriction site followed by the entire 182 predicted sequence for the putative exon 8b and 20-23 bp complementary to either exon 5 or 183 184 exon 7 in order to generate synthetic expression vectors for *Vegfa121b* or *Vegfa165b* (see Table S1). We demonstrated that *Vegfa* amplification by RT-PCR could be obtained when bases 185 "specific for *Vegfaxxxb*" on the primer 5'-end were changed to a GGGGG or an AAAAA stretch 186 [33]. The minimal primer design we defined to be sufficient to obtain artefactual Vegfaxxb-like 187 PCR products perfectly matches the P4 primer described above ([33]; see Figure 2B). Therefore, 188 studies which claimed identification of *Vegfaxxxb* transcripts using primers with \geq 13bp common 189 to Vegfaxxx and putative Vegfaxxb mRNA do not reach required standards of evidence, and 190 their conclusions should be re-evaluated. 191

192

By 2013, the use of the questionable P4 primer design had become widespread in the literature [44,50-52]. This design constitutes a significant departure from initial studies that used the P2 primer with 7bp complementarity to either *Vegfa165* or *Vegfa165b*. The reasons for this drift in primer design are not clear. However, as early as 2006, Bates and colleagues [37] acknowledged that "...RT-PCR is not quantitative and it has not been possible so far to develop isoformspecific qPCR, due to the lack of exon-specific sequences". It seems surprising that primers deemed suitable for specific detection of *Vegfa165b* in standard RT-PCR would not be suitablefor qPCR.

In conclusion, the detection of "*Vegfaxxxb*-like" transcripts could be explained by the use of inadequate RT-PCR methodologies involving 5′-tailing of the primers (also see **BOX1**).

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206 Bioinformatic analysis of alternative splicing: absence of evidence for Vegfaxxxb isoforms.

Using AVISPA, a tool for prediction and analysis of alternative splicing, Barash *et al* reported that "prediction of other splice variations of *Vegfa*, such as the 3' splice site variation in exon 8, are currently not supported by the tool [53] ". Another splice site prediction software ([54]; available at <u>http://www.fruitfly.org/seq_tools/splice.html</u>) identified the canonical acceptor splice site of exon 8 (score of 0.97) but failed to identify the putative exon 8b splice site. Therefore, efficient *in silico* tools do not currently support the existence of the proposed distal splice site in exon 8 of the *Vegfa* gene.

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215 Detection of the *Vegfaxxxb* transcripts: evidence of absence by unbiased RNA-seq analysis.

As detailed above, the most parsimonious explanation for the lack of *Vegfaxxxb* PCR products is that such transcripts do not exist. However, a theoretical possibility remains that failure to detect *Vegfaxxxb* transcripts results from the targeted nature of the approach (also see **BOX2**). If so, the analysis of RNA datasets obtained through the use of an alternative, unbiased methodology, such as RNA-seq, should clarify this matter. Indeed, "RNA-seq represents the method of choice for the discovery of alternative splicing events across tissues [55]".

222

Therefore, we investigated existence of *Vegfaxxxb* transcripts in publicly available RNA-seq datasets generated from multiple human tissues [29]. Our extensive analysis confirmed all known *Vegfa* splice variants and also identified novel transcripts. In particular >40000 *Vegfa* transcripts were uncovered that spanned the junction between exon 7 and exon 8 in >10 different human tissues. None of these reads supported the existence of an exon 8b splice site [29].

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229 We then analysed two independent RNA-seq datasets of hypothalamus / pituitary from castrated rams [57] and ovariectomized, estradiol-implanted ewes [56]. This analysis confirmed that *Vegfa* 230 splice variants are phylogenetically conserved, as transcripts identified in sheep correspond to 231 232 those present in human [33]. Indeed, we also identified several novel ovine Vegfa transcripts that correspond to those identified by Bridgett et al in human [29]. Crucially, we identified 2693 233 *Vegfa* transcripts that cover the junction between exon 5 and exon 8 or between exon 7 and exon 234 235 8. None of the reads supported the existence of an exon 8b splice site, hence the existence of ovine *Vegfaxxxb* transcripts [33]. These findings [33] rule out the proposed role for VEGFAxxxb 236 237 isoforms in the control of ovine seasonal breeding proposed by Castle-Miller et al [58]. We further note that the 43bp-long reverse primer (see Table S1), as provided in the erratum of the 238 aforementioned study [58] to "specifically" amplify Vegfa165b by qPCR, would indeed lead to 239 co-amplification of the classical *Vegfa165* isoform, as its sequence is complementary to a stretch 240 of exon 8 located downstream of the putative distal splice site. This implies that qPCR would 241 yield two amplicons, which makes the primer pair inadequate for this use and leads to questions 242 on the validity of the data published by Castler-Miller et al [58]. In summary, the failure to 243 detect Vegfaxxxb mRNA in human and ovine RNA-seq data strongly suggests that these 244 transcripts do not exist (also see BOX3). 245

246

247 No *Vegfaxxxb* transcripts: detection of VEGFAxxxb proteins is an artefact.

Absence of endogenous *Vegfaxxxb* mRNA implies absence of endogenous VEGFAxxxb proteins. Consequently, studies which investigated VEGFAxxxb levels in tissues or plasma, using anti-VEGFAxxxb antibodies and ELISA kits, or those which relied exclusively on overexpression or injection of recombinant VEGFAxxxb proteins have questionable physiological relevance (see **Table S1**; e.g. [18,59-74]).

253

The first antibody against VEGFAxxxb isoforms (MVRL56/1) was developed by Woolard et al 254 2004 [15] and was raised against the synthetic peptide TCRSLTRKD which corresponds to the 255 nine amino acid C-terminal sequence of human VEGFA165b (see Figure 1C). This peptide is 256 only six amino acids different from the peptide that corresponds to the C-terminal sequence of 257 VEGFA165 (TCRCDKPRR). This antibody was made commercially available and distributed 258 by R&D (MAB3045) and Abcam (Ab149940; [41]). A sandwich ELISA kit that utilizes the 259 260 MAB3045 antibody is also available via R&D (#DY3045) who also supply a recombinant human VEGFA165b protein (#3045-VE-025). Interestingly, Abcam removed Ab149940 from 261 262 their catalog in 2015 as it "did not meet the quality criteria". Specifically, the antibody yielded "low signal-to-noise ratio in immunofluorescence and detected multiple non-specific bands in 263 western-blot" (communication from Abcam). The use of MVRL56/1 for analysis in the same 264 tissue has sometimes led to opposite findings. For instance, using normal breast tissue, Catena et 265 al [20] found no VEGFAxxxb staining while Qiu et al [75] observed strong staining. When used 266 in western-blot studies, MVRL56/1 yields multiple bands (i.e. smear) that span a broad range of 267 molecular weights. These bands have been considered to represent monomers, dimers or large 268 complexes of VEGFAxxxb isoforms [36,37]. However, definitive evidence that these bands 269 correspond to various forms of VEGFAxxxb is still lacking. 270

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As summarized in **Table S1**, virtually all studies aimed at detection of VEGFAxxxb proteins have relied on the use of the MVRL56/1 antibody. Origene Europe also sells a different antibody against hVEGFA165b C-Terminal peptide (the exact sequence of the epitope is not provided; #DM3615P). Finally, an "Anti-VEGFA111b antibody" has been raised [76] using "synthetic peptide fragments of the 8 amino acids CRSLTRKD". Contrary to what the authors claim, this antibody would not be specific for the VEGFA111b isoform, since the epitope corresponds to the 8 amino acid C-terminal sequence shared by all predicted human VEGFAxxxb isoforms.

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Surprisingly, the original anti-human VEGFAxxxb antibody (MVRL56/1) has been used to 280 demonstrate the presence of VEGFAxxxb in multiple species that do not share the same C-281 terminal sequence (see Figure S1). For instance, it has been used in sheep [58] for which the 282 predicted sequence for the last 9 amino acids of the C-Terminal would be TCRCLTRKD, 283 284 therefore slightly divergent from that in human (underlined, Cys instead of Ser). This would reduce the size of the VEGFAxxxb-specific epitope recognized by the anti-VEGFAxxxb 285 antibody to the LTRKD sequence. Having shown that Vegfaxxxb transcripts do not exist, we 286 surmised that VEGFAxxxb-like immunostaining is accounted for by cross-reactivity of the 287 antibody with one or several protein(s) unrelated to VEGFA but bearing an epitope of similar 288 sequence to that of the putative LTRKD sequence of VEGFAxxxb. We searched human protein 289 databases using BLASTP, which led to the unambiguous identification of 10 proteins harboring 290 an LTRKD motif [33]. The MVRL56/1 anti-human VEGFAxxxb antibody was also used to 291 detect endogenous VEGFAxxxb isoforms in tissues from mouse [65,75,77] and rat [45,60], 292 species which, as already noted by us and others ([13,28]; see Figure S1), would present a 293 distinct CRPLTGKTD motif at the C-term of VEGFAxxxb, divergent from the human sequence. 294 Specifically, we found that in mouse, this antibody detects unidentified proteins that can be 295 mistaken for VEGFA isoforms and have raised this as a serious concern [28]. These non-specific 296

proteins were present in mouse cell extracts and their conditioned media as well as mouse tissues. Although the suggestion by Bates *et al* [50] that these findings might be due to artefactual detection of mouse IgG could potentially be correct when testing mouse tissues, this possibility is excluded in cell culture and serum-free conditioned media. Taken together, we conclude that the anti-VEGFAxxxb antibody might cross-react with a large number of endogenous proteins (>10) across different species but none of these belong to the VEGFAxxxb family.

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306 Concluding remarks

We have reviewed current evidence for the proposed existence of a Vegfaxxxb family of 307 transcripts which would yield VEGFAxxxb proteins bearing anti-angiogenic properties. 308 309 Inadequate PCR methodology which likely led to the erroneous identification of such transcripts has been identified and characterized. Multiple published studies, along with our additional 310 analyses of RNA-seq data have failed to identify any Vegfaxxxb-specific splicing events. Thus 311 we conclude that Vegfaxxxb transcripts do not exist in vivo and therefore challenge the view that 312 endogenous VEGFAxxxb proteins have any physiological relevance. Consequently, efforts to 313 develop therapeutics to modulate VEGFA activity should not be based upon modification of a 314 splicing event that is not supported by evidence. The story of *Vegfaxxb* splicing reinforces the 315 importance of supporting new findings using orthogonal techniques prior to basing subsequent 316 studies upon them (see Outstanding Questions). 317

318

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321

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- 498

499 Glossary

- 500 Alternative splicing: process by which splice sites in precursor (pre)-mRNA are differentially
- selected to produce multiple mRNA and protein isoforms. This process diversifies the functional
- 502 characteristics of genes and drastically expands the potential repertoire of protein variants.
- 503 Nearly all multi-exon genes have at least one splice variant.

VEGFA (Vascular Endothelial Growth Factor A): a key member of the family of growth factors
which plays a prominent role in angiogenesis - the growth of blood vessels - both in health and
disease (tumorigenesis and retinopathies).

507 **5'-tailing**: addition of nucleotides on the 5'-end of a PCR primer. These nucleotides are not 508 complementary to the target mRNA but do not impair PCR processivity and aid further cloning 509 of the PCR amplicon.

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- 511
- 512
- 513 **Text BOXES**
- 514 BOX1
- 515 Further technical issues with the use of qRT-PCR

Since primers for standard PCR were not suitable for qPCR, Bates and co-workers developed an 516 alternative method, which we might call "indirect subtractive strategy" [25]. This requires the 517 use of two distinct primer pairs. The first pair consists of "an exon 7b forward primer and a 3'-518 UTR Primer (both pan-VEGF)", the sequences of which were not provided. The second pair of 519 primers consists of "an exon 7a forward primer and a reverse primer specific for exon 8a that do 520 521 not detect Vegfaxxxb isoforms". The authors then assume that subtracting what is found with the second primer pair (Vegfaxxx only) to what is found with the first primer pair (total Vegfa: 522 Vegfaxxx + Vegfaxxb) yields the amount of Vegfaxxb. This is an unusual strategy, which is 523 likely to be flawed for a number of reasons. The primary concern is that these are two different 524 pairs of primers, with different sequences, hence divergent optimal Tm's, which amplify 525 fragments of different lengths and nucleotide composition and therefore have differing 526 efficiencies. The use of qPCR is based on the principle that the number of copies doubles at 527 each cycle. This rule is used as a proxy to determine the initial number of copies. Using the 528

values of the slopes for the calibration curves (see Figure 1 in [25]) for both primer pairs allows 529 calculation of an amplification factor (which should ideally be \sim 2), which informs on the 530 efficiency of the qPCR assay. The efficiency should be ~100%, but values between 85-115% are 531 usually judged acceptable, providing the dissociation curves are good [78]. The calculated 532 efficiency values for the primer pairs used by Varey et al [25] are 83.7% and 71.8%. The 533 meaning of this is twofold. First, neither primer pair is appropriate for use in qPCR. Second, 534 different efficiencies are enough to generate a difference when data obtained with the two primer 535 pairs are compared. A broadly similar technique, but with different primers, was used by the 536 same team in a later study [79]. Other issues with RT-PCR and primer design were also found, 537 538 which are detailed in Supplemental text.

539 **BOX2**

540 Ruling out potential issues with mRNA secondary structure

The idea that PCR amplification of *Vegfa165b* might be difficult due to secondary structure of 541 the mRNA in the 3'-UTR has been invoked [9] to explain why we [31,33] and other authors 542 [13,79,80] failed to detect these transcripts. The existence of an unusually stable secondary 543 structure seems unlikely because most regulatory splicing mechanisms are based on the 544 recognition of short degenerate RNA motifs at the exon/intron boundaries rather than secondary 545 structures [82,83]. Indeed, bioinformatic analysis of the ribonucleotidic sequence does not 546 predict formation of potential hairpins that might inhibit reverse transcription (data not shown). 547 Furthermore, classical Vegfaxxx isoforms are readily amplified using primers spanning the 548 putative alternative splice site (see main text). There is therefore no evidence to support the 549 notion that Vegfaxxxb isoforms, which would lack a 66bp stretch present in Vegfaxxx isoforms, 550 would resist PCR amplification. Furthermore, Vegfa165b transcripts expressed from a 551 recombinant construct were readily detected by RNA-seq [32] and since the library preparation 552 involved both RT and PCR steps, neither are inherently blocked by the transcript secondary 553

structure. Therefore, the inability to detect an abundant mRNA such as *Vegfa165b* at least in normal tissues by classical RT-PCR can not be explained by the secondary structure of the mRNA.

557 **BOX3**

558 Further RNA-seq evidence for absence of *Vegfaxxxb* mRNAs

Recently developed long read RNA sequencing technologies (e.g. PacBio Iso-Seq and Oxford 559 Nanopore Technology) can measure full length mRNA transcripts [85]. Both individual 560 alternative splicing events and the co-occurrence of multiple events in transcripts such as Vegfa 561 are detected. To determine whether data generated with this newly available approach could 562 detect *Vegfaxxb* mRNAs we analysed a publicly available dataset [85]. Full length transcripts 563 encoding all the commonly reported exon 5 and exon 6 isoforms (VEGFA121, VEGFA165, 564 VEGFA189) were present but no evidence of splicing to generate Vegfaxxxb was detected 565 (Supplemental Figure S2). A notable feature apparent from this full length transcript data is the 566 retention of the intron between exons 3 and 4 in almost 20% of transcripts (which is supported 567 by previous EST data). The regulatory potential of such intron retention events has been widely 568 discussed [86,87]. Our recent analysis of RNA-seq data from tumours grown from fibrosarcomas 569 expressing VEGFA188 [88] has also failed to identify reads corresponding to the exon 7-8b 570 splice site, even though endogenous expression of multiple Vegfaxxx isoforms could be detected 571 from the stromal cells (Supplemental Figure S3). To better understand the role of alternative 572 splicing in the regulation of *Vegfa* expression, research should be focused upon such events that 573 are supported by strong experimental evidence. 574

575

576 Figure Legends

577 **Figure 1**: Structure and sequence of the human *Vegfa* gene and VEGFAxxx/VEGFAxxxb 578 isoforms. A/ Schematic of the human *Vegfa* gene locus. The gene is comprised of 8 exons. B/

Schematic of the longest human VEGFA protein (VEGF206, see GenBank accession number 579 NM 001171623). The stop-codon (TGA, in red) is located in exon 8 and the sequence of the last 580 9 amino acids at the C-terminus of the protein is provided below. The location of flanking PCR 581 primers O14I/O16I is also indicated (see panel F). C/ Schematic of the putative human 582 VEGFA206b protein (GenBank accession number NM 001033756). The stop-codon (TGA, in 583 red) is located in exon 8 – renamed exon 8b due to the usage of a distal splice site – and the 584 sequence of the last 9 amino acids at the C-terminus of the protein is provided below. Note that 585 only the sequence of the last 6 amino acids at the C-terminus differ between VEGFA and 586 VEGFAb isoforms. D/ Nucleotide sequence (from nt 1716 to nt 1808) of the 5'-end of human 587 Vegfa exon 8 and deduced amino acid sequences corresponding to the C-term of VEGFAxxx and 588 VEGFAxxxb isoforms; corresponding nucleotide triplets are underlined. The reference sequence 589 used for nucleotide annotation is Genbank NM 001171623. The usual splice site (mauve and 590 591 underlined) leading to VEGFAxxx isoforms (as shown in B) and the putative alternative splice site (green and underlined) leading to VEGFAxxxb isoforms (as shown in C) are shown. The 592 respective stop codons are also shown (red and bold). Also note that exon 8 and exon 8b would 593 share an AT dinucleotide sequence at their 5'-end (grey boxes and bold). E/ Schematic of the 594 four most abundant classical hVEGFA splicing isoforms and of their corresponding sister 595 isoforms of the proposed hVEGFAxxxb family : both families differ only by the alternative use 596 of either exon 8a or 8b, as described in panels B and C using VEGFA206 as an example. 597 Adapted from Bridgett et al [32]. F/ Agarose gel electrophoresis of PCR products obtained using 598 flanking primers O14I/O16I and cDNA from ovine medio-basal hypothalamus (see panel B for 599 location). Gel extraction, cloning and sequencing of the three bands revealed products encoding 600 ovine homologs of human VEGFA206 and VEGFA189 (562 and 544bp; sizes too close to be 601 separated on the gel), VEGFA165 (490bp) and VEGFA121 (358bp). No single VEGFAxxxb 602 PCR product was obtained. For further information, see Lomet et al. [33]. G/ Schematic of the 603

signaling pathway elicited by VEGFA isoforms. Classical VEGFA proteins bind both VEGFR2
homodimer and the Neuropilin1 (NRP1) co-receptor while VEGFAxxxb isoforms fail to bind
NRP1, impairing intracellular transduction pathways.

607

Figure 2: Issues with RT-PCR primer design: how 5'-tailing led to erroneous identification of 608 Vegfaxxxb transcripts. A/ Schematic of the putative junction between exon 7 (blue) and the 609 putative exon 8b (green and black) of Vegfaxxxb transcripts and location of the PCR primers 610 used to identify Vegfaxxb transcripts. Note that exon 7 and the AT dinucleotide (green) at the 611 end of exon 8 sequences would be common to both classical Vegfaxxx and Vegfaxxxb isoforms. 612 613 The number of non-isoform specific nucleotides is provided in blue; note that this number has substantially increased (from 7bp to 17bp) throughout the years (primers P1 through to P7). 614 Sequences in orange represent 5'-tailing. Primers P1-P6 are reverse primers while primer P7 is a 615 616 forward primer, as indicated by the arrows. B/ Minimal sequence requirement to obtain PCR amplification of ovine Vegfaxxxb-like transcripts through 5'-tailing as defined in Lomet et al 617 [33]. The error-prone primer consists of 11 nucleotides in exon 7 and 2-3 nucleotides of exon 8 618 (in green, shared by isoforms), then any nucleotide can be added on the 5'-end of the primer 619 providing the number is sufficient (6 or more) to reach a Tm compatible with the PCR 620 621 methodology (see **Figure 3**). Note that this primer design corresponds to primer P4.

622

Figure 3: Basic principles of PCR and how 5'-tailing is achieved. A/ Principle of standard PCR. After reverse transcription, cDNAs are submitted to 30-40 cycles of denaturation/annealing and extension during which a pair of primers (forward and reverse) allow efficient amplification of the target sequence they flank. B/ To anneal to the cDNA, the melting temperature (Tm) of the primer must be higher than the annealing temperature. C/ The Tm of the primer depends on its nucleotide composition and its length, with long GC-rich primers having higher Tms than short

23

AT-rich primers. A series of 3 efficient primers is shown, the one on the top is the shortest. D/ The shortest efficient reverse primer as defined in C can be flanked on its 5'-end with a large stretch of additional non-specific extra-bases (for instance restriction sites for further cloning purposes), which do not anneal to the target and do not impair PCR efficiency and processivity: this is 5'-tailing. E/ The additional non-specific 5'-tail is added to the amplicon as the PCR progresses. At the end of the 30-40 cycles, all amplicons are flanked with the additional, nonspecific stretch of the reverse primer.

636

637 Supplementary Material

638 Includes Supplemental Text, Supplemental table S1, Supplemental Figues S1-S3

Authors	Journal	Issue:Page	Year	Species/Tissue	PCR	IHC WB ELISA	Recomb. prot Over-exp
Bates et al	Cancer Res	62:4123-4131	2002	Human / Kidney	VEGF165b exon 8b/ 7 - Reverse primer : 5'-TCAGTCTTTCCTGGTGAGAGAGATCTGCA-3'		Home made
					Amplification of full-length VEGF165b with a reverse primer located downstream in the 3'UTR.		
Cui et al	Am J Physiol Renal Physiol	286:F767-773	2004	Human / Kidney	Same primers as Bates <i>et al</i> 2002.		
Woolard <i>et al</i>	Cancer Res	64:7822-7835	2004	Various	Same primers as Bates <i>et al</i> 2002.	Production monoclonal antibody hVEGF165b	Home made
Perrin <i>et al</i>	Diabetologia	48:2422-2427	2005	Human / Eye	Same primers as Bates <i>et al</i> 2002.	Home made ELISA	
Konopatskaya <i>et al</i>	Mol Vis	12:626-632	2006	Mouse / Eye	No		3045-VE-025
Bates et al	Clin Sci (Lond)	110:575-585	2006	Human / Placenta	Same primers as Bates <i>et al</i> 2002.	MAB3045 Home made ELISA	3045-VE-025
Cebe Suarez et al	Cell Mol Life Sci	63:2067-2077	2006	Human cell line (HUVEC)	No	MAB3045	Home made
Ribeiro <i>et al</i>	Mol Reprod Dev	74:163-171	2007	Swine / Ovary	VEGF164b exon 8b/ 7 - Reverse primer : 5'-TCCTGGTGAGAGATCTGCAAG-3' Note : level << VEGF164 (2 ¹² fold)		
Pritchard-Jones <i>et al</i>	Br J Cancer	97:223-230	2007	Human / Melanoma tissue	VEGF165b <u>HindIII RS</u> - exon 8b/ 7 - Reverse primer : 5'-TT <u>AAGCTTTCAGTCTTTCCTGGTGAGACTGCA</u> -3' Note : GAT missing from oligo ; see Bates <i>et al</i> 2002 Invalid primer.	MAB3045	
Schumacher <i>et al</i>	J Am Soc Nephrol	18:719-729	2007	Human / Kidney	VEGF both isoforms <u>BamHI RS</u> – 3'UTR Reverse primer : 5'-AT <u>GGATCCGTATCAGTCTTTCCT</u> -3' Note : short primer, theoretical fusion T° = 41°C		
Ergorul <i>et al</i>	Mol Vision	14:1517-1524	2008	Rat / Retina	No	Ab14994	
Qiu et al	Faseb J	22:1104-1112	2008	Mouse / Various tissues	Primers use to « detect specifically the transgene » Forward : 5'-TCAGCGCAGCTACTGCCATC-3' Reverse : 5'-GTGCTGGCCTTGGTGAGGTT-3' Note : Forward is within exon3 and Reverse is within exon → actually detect all Vegfa.	MAB3045	Yes – TG mice

					Invalid primers \rightarrow Invalid screening of TG mice.		
Bevan <i>et al</i>	Nephron Physiol	110:57-67	2008	Human / Kidney	No	MAB3045	
Kawamura <i>et al</i>	Cancer Res	68:4683-4692	2008	Cell lines	No		Home made
Varey <i>et al</i>	Br J Cancer	98:1366-1379	2008	Human / Colon	Original indirect subtractive strategy : Use of « An exon 7b forward primer and a 3'UTR Primer (both pan-VEGF) » - Sequences not provided Then use of « an exon 7a forward primer and a reverse primer specific for exon 8a that did not detect Vegfxxxb isoforms ». Then subtraction would yield Vegfxxxb. Invalid method.	MAB3045	Yes
Diaz <i>et al</i>	Int J Cancer	123 :1060-1067	2008	Human / Colon	Same primers as Bates et al 2002.	MRVL56/1	
Nowak et al	J Cell Sci	121:3487-3495	2008	Cell lines	VEGF both isoforms <u>BamHI RS</u> – 3'UTR 5'-AT <u>GGATCCGTATCAGTCTTTCCTGG</u> -3'	MAB3045 & clone 264610/1	Home made
Rennel <i>et al</i>	Br J Cancer	98:1250-1257	2008	Human / Prostate Mouse	VEGF165b <u>HindIII RS</u> - exon 8b/ 7 5'-TT <u>AAGCTTTCAGTCTTTCCTGGTGAGAGATCTGCA</u> -3'	Home made ELISA	Home made
Rennel <i>et al</i>	Eur J Cancer	44:1883-1894	2008	Cell lines Mouse	No	MAB3045	Home made
Miller-Kasprzak and Jagodinski	Biomed Pharmacother	62:158-163	2008	Human / Lung cell line	VEGF121b exon 5/8b - Forward primer : 5'-GAAAATCTCTCACCAGGAAA-3' Note : one A missing from oligo. VEGF165b exon 8b/ 7 - Reverse primer : 5'-GTGAGAGATCTGCAAGTACG-3'	MAB3045	
Artac et al	Biol Reprod	81:978-988	2009	Rat / Ovary	VEGF165b exon 8b/ 7 - Reverse primer : 5'-GGTGAGAGGTCTGCAAGTACGTT-3'	MAB3045	
Bills et al	Clin Sci (Lond)	116:265-272	2009	Human / Plasma	No	MAB3045 Home made ELISA	
Rennel <i>et al</i>	Br J Cancer	101:1183-1193	2009	Human / Colon	VEGF121b exon 5/8b - Forward primer : 5'-GAAAAATCTCTCACCAGGAAA-3'	MAB3045	DY3045
Baltes-Breitwisch et al	Reproduction	140:319-329	2010	Rat / Testes	Same primers as Artac et al. 2009.	Ab14994	
Catena <i>et al</i>	Molecular Cancer	9 :320	2010	Human cell lines	VEGF121b – <u>HindIII RS</u> – exon 8b/ exon5 Reverse primer : 5'-TT <u>AAGCTT</u> TCAGTCTTTCCTGGTGAGAGAGATTTTTCTT GTCTTGCTCTATC - 3' VEGF165b – <u>HindIII RS</u> – exon 8b/ exon7 Reverse primer : 5'-TT <u>AAGCTT</u> TCAGTCTTTCCTGGTGAGAGATCTGCAA GTACGTTCGTTTAACTC - 3'	MAB3045	Yes

Hua <i>et al</i>	Invest Ophtalmol Vis Sci	51:4282-4288	2010	Human / Eye	No	MAB3045	
Magnussen <i>et al</i>	Invest Ophtalmol Vis Sci	51:4273-4281	2010	Mouse / Eye	No	MAB3045	3045-VE-025
Nowak <i>et al</i>	J Biol Chem	285:5532-5540	2010	Cell lines	Same as Nowak <i>et al</i> 2008	MAB3045	
Peiris-Pagès <i>et al</i>	J.Pathol	222:138-147	2010	Cell lines	Same as Nowak et al 2008	MAB3045	DY3045 3045-VE-025
Qiu et al	J Am Soc Nephrol	21:1498-1509	2010	Cell lines Mouse	Same invalid primers as Qiu et al 2008 to « detect specifically the transgene » → Invalid screening of TG mice.	Home made ELISA & clone 264610/1	TG-mice DY3045
Merdzhanova <i>et al</i>	Oncogene	29: 5392-5403	2010	Mouse & Human Cell lines	VEGF165b exon 8b/ 7 - Reverse primer : 5'-TGGTGAGAGATCTGCAAGTACGTT-3'		
Zhao <i>et al</i>	Exp Eye Res	93:921-926	2011	Mouse / Retina	No	Ab14994	
Amin <i>et al</i>	Cancer Cell	20:768-780	2011	Cell lines	Same primer as Nowak et al 2008.		
Xu et al	Gene	487:143-150	2011	Human-Cat-Rabbit- Rat/ Various tissues	Human : VEGF165b exon 8b/ 7 Reverse primer : 5'-TGGTGAGAGATCTGCAAGTACGTT-3' Cat : VEGF165b exon 8b/ 7 Reverse primer : 5'-TGGTGAGAGGTCTGCAAGTACGTT-3' Rabbit : VEGF165b exon 8b/ 7 Reverse primer : 5'-CGGTGAGAGGTCTGCAAGTACGTT-3' Rat : VEGF165b exon 8b/ 7 Reverse primer : 5'-GGTGAGAGGTCTGCAAGTACGTT-3'		
Manetti <i>et al</i>	Circ Res	109:e14-26	2011	Human / Skin	Same primer as Rennel et al. 2008	Ab14994	
Baba <i>et al</i>	Dev Dyn	241:595-607	2012	Human / Eye	VEGF165b exon 8b/ 7 - Reverse primer : 5'-TCCTGGTGAGAGA <u>ATCTGCAA</u> -3'	Ab14994	
Oltean <i>et al</i>	Am J Physiol Renal Physiol	303:F1026-1036	2012	Mouse / Kidney	Same invalid primers as Qiu et al 2008 to « detect specifically the transgene » → Invalid screening of TG mice.		TG-mice
Qiu <i>et al</i>	Reproduction	143:501-511	2012	Marmoset / Ovary Mouse	No	Home made ELISA & clone 264610/1	
Clifford et al	J Immunol	189:819-831	2012	Human / Cell lines	VEGF165b exon 8b/ 7 - Reverse primer : 5'-AGAGAGATCTGCAAGTACGTTCG-3' VEGF189b exon 8b/ 7 - Reverse primer : 5'-GTGAGAGATCTGCAAGTACG-3'		
					5 Grandra rider rider rider rider		

					5'-GTGAGAGATCTGCAAGTACG-3'		
Beazley-Long et al	Am J Pathol	183:918-929	2013	Human / Brain Rat / Brain	No	MAB3045	3045-VE-025
Gu et al	Biochem Biophys Res Comm	441:18-24	2013	Human / Cell lines	VEGF111b – « Reverse primer » : 5' – AATGCAGATGTGACAAGCCGAG – 3' Actually a forward primer at junction exon4/8a. Invalid → Identification of Vegf111b invalid. VEGF165b <u>HindIII RS</u> - exon 8b/ 7 - Reverse primer : 5'-TT <u>AAGCTT</u> TCAGTCTTTCCTGGTGAGAGATCTGCA-3'	MRVL56/1 & Home Made αVEGF111b	
Delcombel <i>et al</i>	Angiogenesis	16:353-371	2013	HEK293 cells	VEGF111b exon8b/4- Reverse primer : 5'-TCCTGGTGAGAGATCTGCATTCAC-3' VEGF121b exon8b/5- Reverse primer : 5'-GTCTTTCCTGGTGAGAGAGATTTTCTT-3' Note : primer issue → AT missing between exon8b and exon5. Invalid. VEGF165b exon8b - Reverse primer : 5'-CGATCGTTCTGTATCAGTCTTTCCT-3' Note : internal to exon 8 → PCR product = classical mVEG Invalid.		
Hulse <i>et al</i>	Neurobiol Dis	71:245-259	2014		Strategy similar to Varey et al 2008 but different Primers ; « primers specific for VEGF-A165a » are: Forward primer exon 7: 5'-GTTCAGAGCGGAGAAAGCAT-3' Reverse primer exon 8a : 5'- TCACATCTGCAAGTACGTTCG-3' → This primer covers exon8a/7	Ab14994	DY3045
Kikuchi <i>et al</i>	Nat Med	20:1464-1471	2014	Human	hVEGF165b exon 8b/7 - Reverse primer : 5'-GTGAGAGATCTGCAAGTACG-3' 		3045-VE-025
Hulse <i>et al</i>	Clin Sci	129:741-756	2015	Cell culture Rat	No		3045-VE-025
Oltean <i>et al</i>	J Am Soc Nephrol	26:1889-1904	2015	Human / Kidney	VEGF165b exon 8b/7 - Reverse primer : 5'-GTGAGAGATCTGCAAGTACG-3'	DY3045	
Vencappa <i>et al</i>	Am J Transl Res	7:1032-1044	2015	Cell culture Mouse	No	Ab14994	3045-VE-025
Li et al	J Translat Med	13 :164	2015	Human cell lines	Same primers as Gu et al 2013	Same as	Yes

						Gu <i>et al</i> 2013	
Hulse <i>et al</i>	Neurobiol Dis	96:186-200	2016	Rat	No	Ab14994	
Castle-Miller <i>et al</i>	Proc Natl Acad Sci USA	114:E2517-2523	2017	Sheep / Pituitary	Multiple issues with primers leading to an erratum. VEGFAxxxb-specific reverse primer : 5'-CGGCGGCTATGGGTCGTTCTGTGTCAGTCTTTCCT GGTGAGAC-3' 43bp. Unusual. Targets a sequence downstream of putative distal splice : located in exon 8. Non-isoform specific. → qRT-PCR data invalid.	Home made Clone56/8 & DY3045	
Ved <i>et al</i>	Clin Sci (Lond)	131:1225-1243	2017	Rat / Retina	No	Home made Clone56/8	Yes
Pruszko <i>et al</i>	EMBO reports	18:1331-1351	2017	Human cell lines	VEGF121b exon 8b/5 - Reverse primer : 5'-CTTTCCTGGTGAGAGATTTTTCTTGTC-3' VEGF165b exon 8b/7 - Reverse primer : 5'-CCTGGTGAGAGATCTGCAAGTAC-3'	Ab14994	
Hueso <i>et al</i>	Scientific Reports	7 :9962	2017	Human Serum / Heart	No	MAB3045	MBS109074
Chesnokov <i>et al</i>	PeerJ	6:e4915	2018	Human / Tissues	VEGFxxxb exon 7/8b - Forward primer : 5'-ACGTACTTGCAGATCTCTCACCA-3' Underlined bases correspond to either end of exon7 OR exon 8.		
Blochowiak et al	Adv Clin Exp Med	27:83-90	2018	Human / Parotid	No	DM3615P	
Boudria <i>et al</i>	Oncogene	AOP		Mice / Cell lines	No	Yes	Yes
Stevens et al	Nephron	139:51-62	2018	Mice	No	No	Yes / TG mice

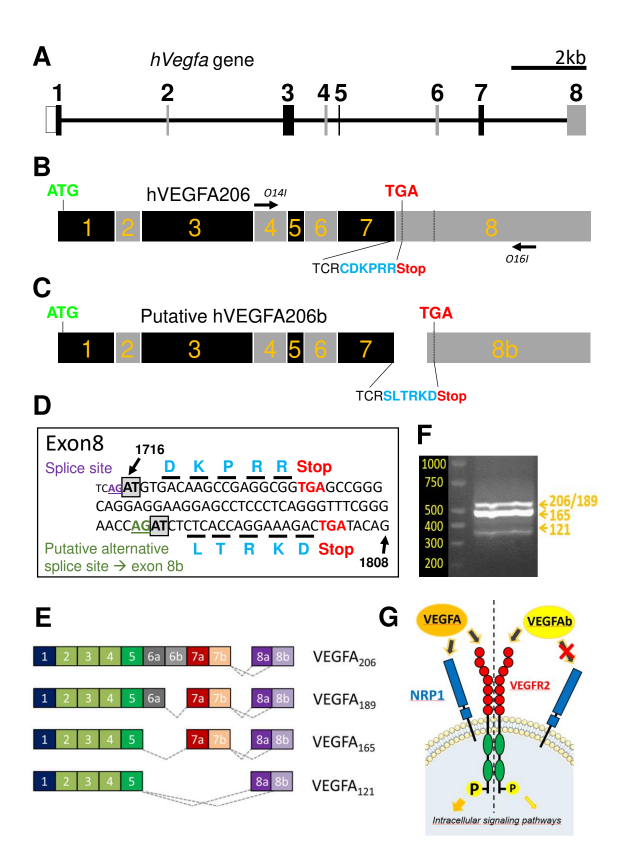
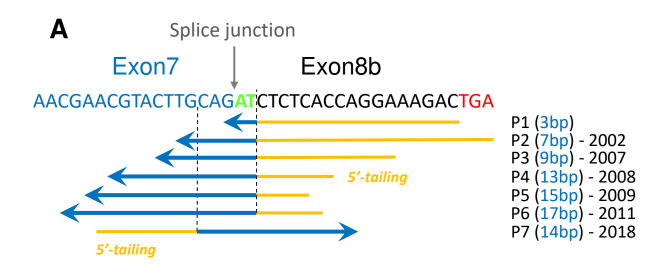
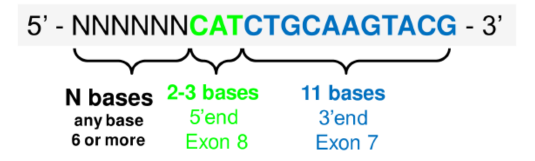


Figure 1



Β



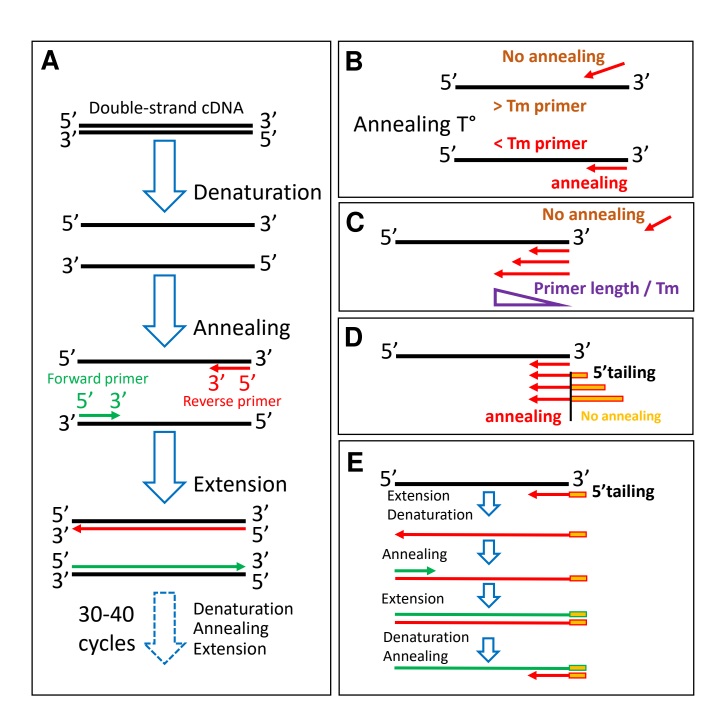


Figure 3

Supplemental Text

Debunking the myth of the endogenous anti-angiogenic *Vegfaxxxb* transcripts. Hugues Dardente^{1*}, William R English², Manoj K Valluru², Chryso Kanthou² & David Simpson³ ¹PRC, INRA, CNRS, IFCE, Université de Tours, 37380 Nouzilly, France ²Department of Oncology and Metabolism, Tumour Microcirculation Group, University of Sheffield, School of Medicine, Beech Hill Road, Sheffield, S10 2RX, United Kingdom ³Centre for Experimental Medicine, Queen's University Belfast, Belfast, BT7 1NN, Northern Ireland

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Other methodological issues with RT-PCR

We noted further inconsistencies in the literature and not-readily-explainable RT-PCR findings. First, to identify *Vegfa165b*, Bates *et al* [1] used primers designed to detect *Vegfa148*, a transcript identified from kidney by the same team [2] and recently shown to be enriched in some tissues, including kidney [3]. The pair of primers used by Bates *et al* [1] consisted of a forward primer located within exon 7a and a reverse primer located in the 3'-UTR, downstream of the putative alternative 8b splice site (i.e. flanking primers; **Table S1**). This should yield at least two PCR products of 185bp and 151bp, corresponding to *Vegfa165* and *Vegfa148*. Should the *Vegfa165b* transcript exist, one would expect an extra PCR product at 120bp. Hence, 3 bands should be discernible on the gel, with the one corresponding to *Vegfa148* located in between those for *Vegfa165* and *Vegfa165b*. This was not the case. This is puzzling considering these primers were in the first instance "designed to detect *Vegfa148*". The same issue applies to the study by Schumacher *et al* [4] who used the same primers. Furthermore, *Vegfaxxb* transcripts were not identified by

Whittle *et al* [2], who used a pair of *ad hoc* primers for this (forward primer in exon 7a and reverse primer downstream of the putative 8b splice site).

In follow-up studies from the team led by Bates and Harper, DNA rulers were often missing and sequencing of the PCR products was not reported [e.g.; 5-8], and therefore the identity of PCR products could not be confirmed. Rather, to validate the existence of an entire family, the authors used assertions such as "size compatible with *Vegfa165b*" [9], "bands corresponding to VEGFA1xxb isoforms (VEGFA165b, VEGFA189b, etc.) were seen ..." [6] but unfortunately not shown, or "multiple isoforms were detected, with bands seen consistent with VEGFA121b (108 bp), VEGFA145b (219 bp) and VEGFA165b (240 bp)". In particular, Cui *et al* [9] used isoform-specific reverse primers to amplify either *Vegfa165* or *Vegfa165b*. These primers were used in combination with the same forward primer, located in exon 4. Therefore RT-PCR for *Vegfa165* is expected to yield at least 3 major bands. Considering the "sister family of *Vegfa189b* and *Vegfa165b*). However, only products of "sizes compatible with" the presence of *Vegfa189* and *Vegfa165b* were evident on the gel (see [9]; their Fig.3).

Finally, primers used to "detect specifically the transgene" [10,11] in a transgenic mouse model, which over-expresses *Vegfa165b* in the mammary gland, are located within exon3 (forward primer) and exon4 (reverse primer). Therefore, this primer pair will detect all *Vegfa* isoforms (see **Table S1**), which casts doubts on this mouse model.

¹ Bates, D.O. et al. (2002) VEGF165b, an inhibitory splice variant of vascular endothelial growth factor, is down-regulated in renal cell carcinoma. Cancer Res 62, 4123-4131

² Whittle, C. *et al.* (1999) Heterogeneous vascular endothelial growth factor (VEGF) isoform mRNA and receptor mRNA expression in human glomeruli, and the identification of VEGF148 mRNA, a novel truncated splice variant. *Clin Sci (Lond)* 97, 303-312

3 Bridgett, S. *et al.* (2017) RNA-Sequencing data supports the existence of novel VEGFA splicing events but not of VEGFAxxxb isoforms. *Sci Rep* 7, 58-017-00100-3

4 Schumacher, V.A. *et al.* (2007) Impaired glomerular maturation and lack of VEGF165b in Denys-Drash syndrome. *J* Am Soc Nephrol 18, 719-729

5 Woolard, J. et al. (2004) VEGF165b, an inhibitory vascular endothelial growth factor splice variant: mechanism of action, in vivo effect on angiogenesis and endogenous protein expression. *Cancer Res* 64, 7822-7835

6 Perrin, R.M. *et al.* (2005) Diabetic retinopathy is associated with a switch in splicing from anti- to pro-angiogenic isoforms of vascular endothelial growth factor. *Diabetologia* 48, 2422-2427

7 Bates, D.O. *et al.* (2006) The endogenous anti-angiogenic family of splice variants of VEGF, VEGFxxxb, are down-regulated in pre-eclamptic placentae at term. *Clin Sci (Lond)* 110, 575-585

8 Peiris-Pages, M. *et al.* (2010) Balance of pro- versus anti-angiogenic splice isoforms of vascular endothelial growth factor as a regulator of neuroblastoma growth. *J Pathol* 222, 138-147

9 Cui, T.G. *et al.* (2004) Differentiated human podocytes endogenously express an inhibitory isoform of vascular endothelial growth factor (VEGF165b) mRNA and protein. *Am J Physiol Renal Physiol* 286, F767-73

10 Qiu, Y. *et al.* (2008) Mammary alveolar development during lactation is inhibited by the endogenous antiangiogenic growth factor isoform, VEGF165b. *Faseb j* 22, 1104-1112

11 Qiu, Y. et al. (2010) Overexpression of VEGF165b in podocytes reduces glomerular permeability. J Am Soc Nephrol 21, 1498-1509

Antibodies and ELISA kits (see text and Supplemental Table1):

MAB3045 from R&D : Monoclonal Mouse IgG_1 Clone # 56-1; directed against human VEGF_{165b} synthetic peptideTCRSLTRKD.

https://www.rndsystems.com/products/human-vegf-165b-antibody-56-1_mab3045

Note : another Ab (raised against the same 9 amino acids) from clone #264610/1 was used in Nowak et al 2008 – no further information could be found.

Note : for Castle-Miller *et al* 2017 and Ved *et al* 2017 another clone (56/8) was used. No validation provided.

- DM3615P from Origene : Monoclonal Mouse IgG1 Clone #7F17; directed against human recombinant Human VEGF-165b C-terminal peptide
- https://www.acris-anticorps.fr/antibodies/primary-antibodies/vegf165b-dm3615p.htm
- DY3045 from R&D : human VEGF_{165b} Solid Phase Sandwich ELISA <u>https://www.rndsystems.com/products/human-vegf-165b-duoset-elisa_dy3045#product-details</u>
- 3045-VE-025 : Recombinant human VEGFA165b protein <u>https://www.rndsystems.com/products/recombinant-human-vegf-165b-protein_3045-ve</u>
- Ab14994 (clone MRVL56/1) from Abcam : mouse monoclonal antibody discontinued since 2015. https://www.abcam.com/vegf165b-antibody-mrvl561-ab14994.html
- MBS109074 form MyBioSource : hVEGFA165b ELISA kit <u>https://www.mybiosource.com/prods/ELISA-Kit/Human/Vascular-Endothelial-Growth-Factor-</u> 165B-VEGF-165B/VEGF-165B/datasheet.php?products_id=109074

List of additional references only cited in Supplemental Table1 :

1 Amin, E.M. et al. (2011) WT1 mutants reveal SRPK1 to be a downstream angiogenesis target by altering VEGF splicing. Cancer Cell 20, 768-780

2 Baltes-Breitwisch, M.M. *et al.* (2010) Neutralization of vascular endothelial growth factor antiangiogenic isoforms or administration of proangiogenic isoforms stimulates vascular development in the rat testis. *Reproduction* 140, 319-329

3 Hueso, L. *et al.* (2017) Dynamics and implications of circulating anti-angiogenic VEGF-A165b isoform in patients with ST-elevation myocardial infarction. *Sci Rep* 7, 9962-017-10505-9

4 Li, X. *et al.* (2015) VEGF111b, a C-terminal splice variant of VEGF-A and induced by mitomycin C, inhibits ovarian cancer growth. *J Transl Med* 13, 164-015-0522-0

5 Nowak, D.G. *et al.* (2008) Expression of pro- and anti-angiogenic isoforms of VEGF is differentially regulated by splicing and growth factors. *J Cell Sci* 121, 3487-3495

6 Nowak, D.G. *et al.* (2010) Regulation of vascular endothelial growth factor (VEGF) splicing from pro-angiogenic to anti-angiogenic isoforms: a novel therapeutic strategy for angiogenesis. *J Biol Chem* 285, 5532-5540

7 Qiu, Y. et al. (2010) Overexpression of VEGF165b in podocytes reduces glomerular permeability. J Am Soc Nephrol 21, 1498-1509

8 Stevens, M. *et al.* (2018) Vascular Endothelial Growth Factor-A165b Restores Normal Glomerular Water Permeability in a Diphtheria-Toxin Mouse Model of Glomerular Injury. *Nephron* 139, 51-62

		Exon8	Exon8b
			¥
Chinese hamster	<i>GCAG</i> ATGTGACAAGCCAAGGCGG <mark>TC</mark>	AGCCAGGCTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTGTCACCAG-AAAGACCGAC <mark>TA</mark> <mark>A</mark> CCACAC
Vole	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> G	<mark>;A</mark> GCCAGGCTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC <mark>AG</mark>	CCCTCTCACCAT-GTCACCACCACAC
Mouse	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> (<mark>;A</mark> GCCAGGCTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC <mark>AG</mark>	ACCTCTCACCGG-AAAGACCGAT <mark>TAA</mark> CCATGTCACCACCACGC
Rat	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> (IAGCCAGGCTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACTAG	ACCTCTCACCGG-AAAGACCGAT <mark>TAA</mark> CCATGTCACCACCACAC
Deer mice	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> (IAGCCAGGCTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAG-AAAGACCGAC <mark>TAA</mark> CCATGTCACCACCACAC
Mole rat	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> (<mark>;A</mark> GCCAGGCAGAAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAG-AAAGACCGAC <mark>TAA</mark> CTATGTCGCCGCCACCACAC
Beaver	GCAGATGTGACAAGCCAAGGCGG <mark>T</mark> (IAGCCAGGTTGCAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAGGAAAGAT <mark>TGA</mark> TCCAGCACGAC-CGACTACTACAC
Alpaca	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark>	IAGCCGGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ATGTCTCACCAGGAAAGAC <mark>TGA</mark> CACGACTGACAGCCGCTGCCACCA
Goat	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> (AGCCGGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACGTCTCACCAGGAAAGAC <mark>TGA</mark> CACAGAACGACCCATAGCCGCCGC
Sheep	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> G	I <mark>A</mark> GCCGGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ATGT <u>CTC</u> ACC <u>AGG</u> AAA <u>GAC</u> TGACACAGAACGACCCATAGCCGCCGC
Cow	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> (IAGCCGGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACGTCTCACCAGGAAAGAC <mark>TGA</mark> CACAGAACTACCCATAGCCGCCGC
Baboon	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark>	IA <mark>GCCGGGCAGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC</mark> AG	ATCTCTCACCAGGAAAGAC <mark>TGA</mark> TACAGAACGATCGATACAGAAACCACGC
Macaque	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> (IA <mark>GCCGGGCAGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC</mark> AG	ATCTCTCACCAGGAAAGAC <mark>TGA</mark> TACAGAACGATCGATACAGAAACCACGC
Snub-nosed monkey	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> (IN <mark>GCCAGGCAGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC</mark> AG	ATCTCTCACCAGGAAAGAC <mark>TGA</mark> TACAGAACGATCGATACAGAAACCACGC
Human	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> (IAGCCGGGCAGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ATCTCTCACCAGGAAAGAC <mark>TGA</mark> TACAGAACGATCGATACAGAAACCACGC
Coquerel's sifaka	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark> G	J <mark>A</mark> GCCGGGCCGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAGGAAAGACCGACACAGAAGCCACGC
Donkey	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark>	HAGCCAGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCAAGGAAAGAC <mark>TGA</mark> TAGGGAACGACCGAAACAGAAACCACGC
Mouse lemur	GCAGATGTGACAAGCC <mark>G</mark> AGGCGG <mark>T</mark> (HAGCCGGGCAGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAGGAAAGACCGACACAGAACGACCGACACAGAAGCCACGC
Bat	GCAGATGTGACAAGCCAAGGCGGT(HAGCCAGGCTGGAGGAAGGAACCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAGGAAAGATGGATACAGAATGACTGATACAGAAACCACGC
Bat2	GCAGATGTGACAAGCCGAGGCGG <mark>T</mark>	IA <mark>GCCAGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC</mark> AG	ACCTCTCACCAGGAGAGATGGATACAGAATGACTGATACAGAAACCACGC
Star-nosed mole	GCAGATGTGACAAGCCCAGGCGGTG	IA <mark>GCCAGGCTGGAGGAAGGAGCCTCCCTCAGGGTTTCGGGAACC</mark> AG	
Cat	GCAGATGTGACAAGCC <mark>C</mark> AGGCGG <mark>T</mark> ************************************	CCCGGGCTGGAAGAAGGAGCCTCCCTCAGGGTTTCGGGAACCAG	ACCTCTCACCAGGAAAGAC <mark>GA</mark> TACAGAACGACTGACACAGAAACCACGC * *** *

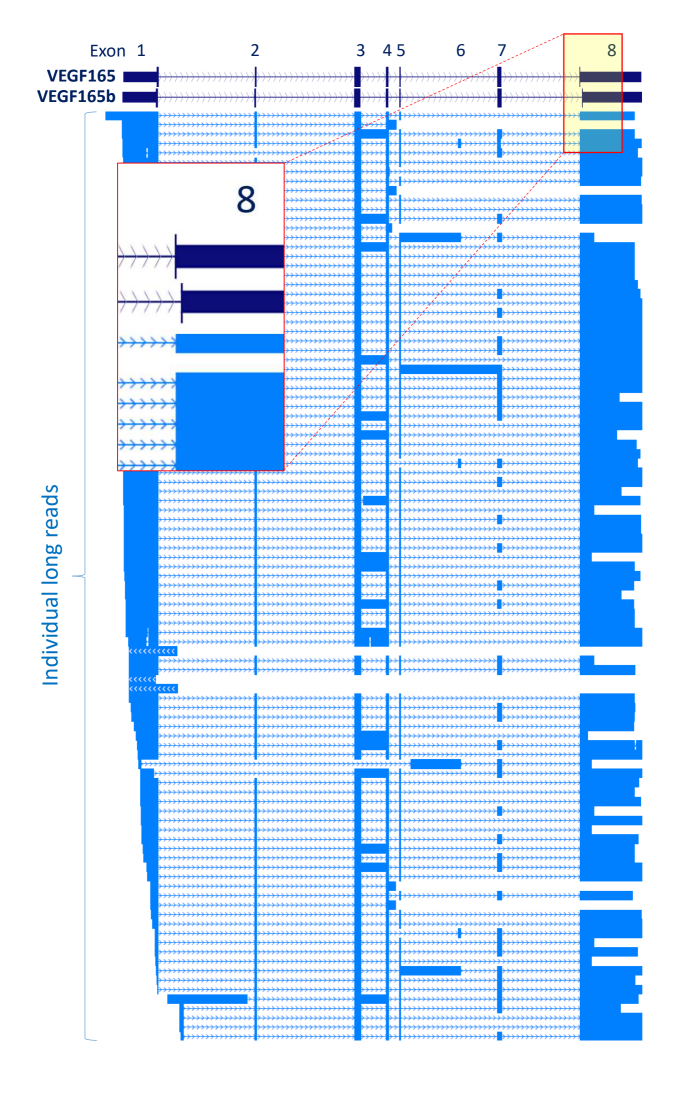
Canonical splice site

Alternative splice site

CCA/CCG/CCC/CCT = Proline.	
22 seq, all identical = CDKPRR	

Chinese hamster	PVTRKTD Putative VEGFAxxxb sequence	000
Deer mice/Mole rat		663
Mouse/Rat	PLTGKTD	
Star-nosed mole	PLTGKD	
Sheep/Alpaca	CLTRKD	
Goat/Cow	RLTRKD	
Human/Babbon/Macaque	SLTRKD	
Beaver/Cat	PLTRKD	
Vole	PLTMSPPHRHRHHRQNSP	
Bat	PLTRKDGYRMTDTETTPLPPHHHH	
Bat2	PLTRRDGYRMTDTETTPLPPHHHHHHRQNNP	
Mouse	PLTRKDRHRTTDTEATPLPPHHRHHRQNRP	
Donkey	PLKERLIGNDRNRNHAAAATTSPSTEQSLIQKPEMKEEETAQSTLGPEGETPAEAFPGG	
Coquerel's sifaka	eq:pltrkdrhrshaaattppspsteqsldqkpemkeeetlrralwvrrarlrqkhsragdqarslleideftemkeeetlrralwvrrarlrqkhsragdqarslleideftemkeestlrkqqarslleideftemkeestlrralwvrrarlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeestlrqkhsragdqarslleideftemkeest	LDSPFRFSCC
	:	

Figure S1



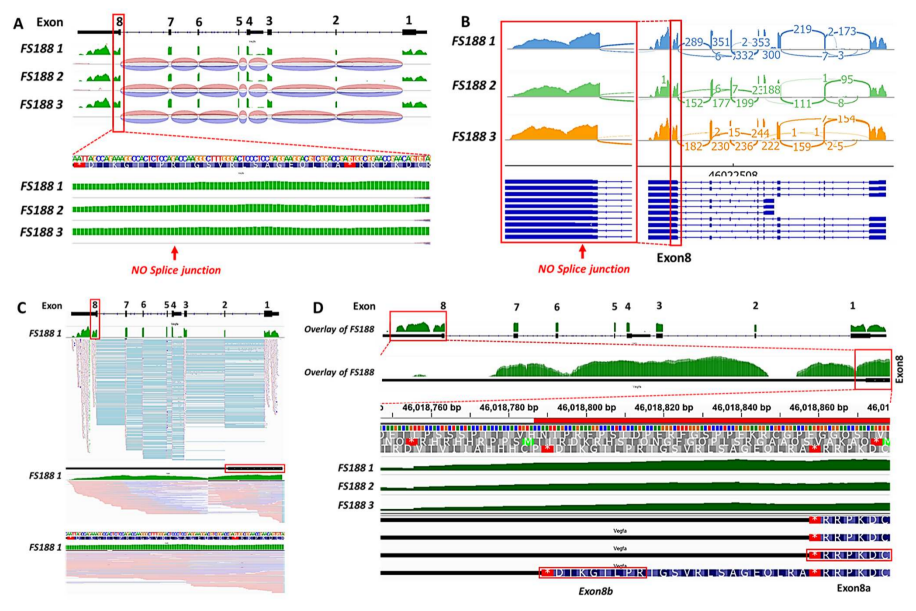


Figure S3