

# 1 **Template plasmid integration in germline genome-edited cattle.**

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## 7 **Author contributions:**

8 A.L.N. and S.S.L. developed the strategy; A.L.N. conducted the experiments; A.L.N., S.S.L.,  
9 and D.A.T. analyzed the data; and A.L.N, S.S.L, K.J.G., D.A.T., M.F.M., and H.L. wrote the  
10 manuscript.

## 11 **Competing Financial Interests:**

12 The authors declare no competing financial interests.

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19 **Abstract**

20 We analyzed publicly available whole genome sequencing data from cattle which were  
21 germline genome-edited to introduce polledness. Our analysis discovered the unintended  
22 heterozygous integration of the plasmid and a second copy of the repair template sequence,  
23 at the target site. Our finding underscores the importance of employing screening methods  
24 suited to reliably detect the unintended integration of plasmids and multiple template  
25 copies.

26 **Article**

27 As genome editing technology evolves, so does our understanding of the unintended  
28 alterations it produces, both in form and frequency. Several sequencing-based methods  
29 have been developed to screen for off-target errors (GUIDE-Seq<sup>1</sup>, SITE-Seq<sup>2</sup>, CIRCLE-Seq<sup>3</sup>,  
30 DISCOVER-Seq<sup>4</sup>), and long-read sequencing of the target site can be used to detect on-  
31 target errors. Each screening approach carries assumptions and biases that may allow  
32 alterations of unexpected types to go undetected. Recent examples of previously  
33 unexpected alterations are complex genomic rearrangements at or near the target site in  
34 mammalian genome editing experiments<sup>5, 6</sup>. The complex rearrangements included  
35 insertions, deletions, inversions, and translocations that were difficult to detect by standard  
36 PCR and DNA sequencing methods.

37 In this study, we analyzed the target site of publicly-available whole genome sequencing  
38 data<sup>7</sup> from genome-edited calves to confirm the intended edit and to screen for potentially  
39 undetected on-target errors (**Supplementary Methods**). The calves were genome-edited<sup>7</sup>,  
40 using transcription activator-like effector nucleases (TALENs) and a repair template for  
41 homology-directed repair (HDR)<sup>8</sup>, to introduce the Celtic polled allele ( $P_c$ ), a variant that  
42 produces the hornlessness (polled) trait in cattle. The  $P_c$  variant, common in some cattle  
43 breeds<sup>9</sup>, is a 212-bp duplication in place of a 10-bp sequence in an intergenic region on  
44 chromosome 1 (chr1:2,429,000-2,429,500; bosTau9). The variant follows an autosomal  
45 dominant inheritance, but the mechanism underlying the association with polled trait is  
46 unknown.

47 Given that the repair template plasmid was delivered in the pCR2.1-TOPO plasmid (**Fig.**  
48 **1a**), we included the plasmid backbone sequence in our comparison of the sequencing reads  
49 with the bovine reference genome. In our analysis, we discovered the presence of the full-  
50 length plasmid backbone in both genome-edited calves (**Supplemental Fig. 1**). While one  
51 allele contained the intended edit, identical to the naturally-occurring  $P_c$  variant (**Fig. 1b**),

52 close inspection revealed integration of the plasmid and a second copy of the repair  
53 template sequence at the target site in the other allele in both calves (**Fig. 1c**). The  
54 plasmid-containing allele (denoted  $P_c^*$ ) was found to have been inserted continuous with  
55 the template ends, producing a duplication of the template and two novel bovine-plasmid  
56 junctions (**Supplemental Fig. 2**). No off-target insertions of the plasmid or the template  
57 were detected.

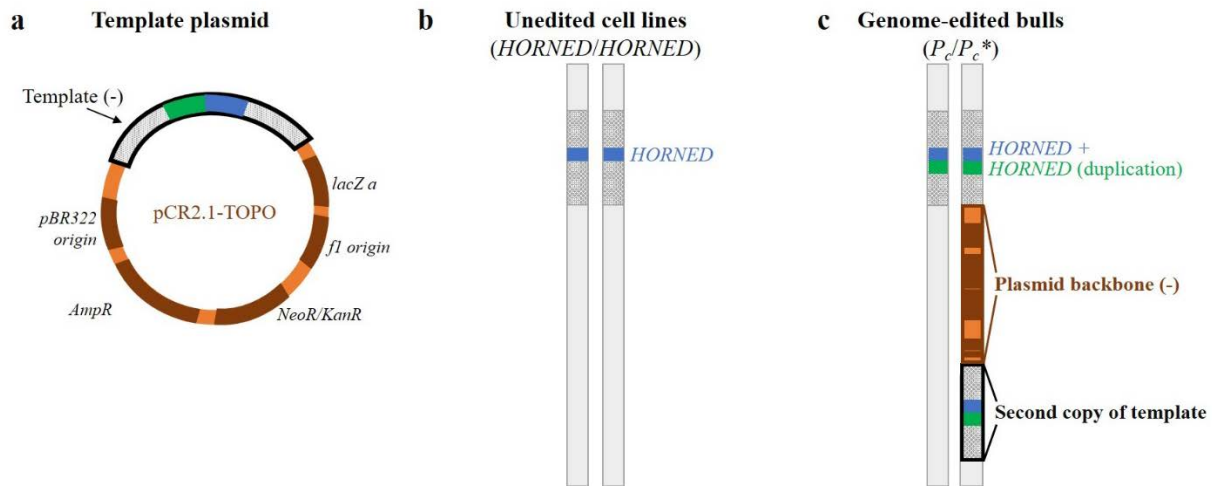
58 Previously, the template plasmid integration was not detected<sup>7</sup>. Probable reasons include:  
59 the plasmid backbone was not included in the sequence alignment, elevated noise at the  
60 target locus, limited signal of the sequencing data, and PCR conditions insensitive to detect  
61 the integrations. The noise was elevated due to the complex sequence context that  
62 obscured the integration: (1) the  $P_c$  variant itself is a duplication of the reference sequence  
63 (*HORNED* allele) in place of a 10-bp sequence, and (2) the target locus is highly repetitive,  
64 potentially masking rearrangements. The signal is limited by the sequencing depth of 20  
65 reads for each DNA base, on average. In an ideal scenario, heterozygosity would result in  
66 10 reads identifying the plasmid insertion but given that the plasmid DNA sequence is not in  
67 the reference genome, the plasmid reads would remain unmapped. The template plasmid  
68 integration was not detected by PCR genotyping<sup>7</sup> due to the following: (1) the expected PCR  
69 amplicons, correctly sized for the  $P_c$  variant (212-bp duplication in place of a 10-bp  
70 sequence), were produced, (2) the primers were not designed to amplify the plasmid, (3)  
71 the amplicons produced by the template plasmid integration were prohibitively large, and  
72 (4) the qualitative nature of the assay was insensitive to the increased number of template  
73 copies (**Supplementary Methods, Supplemental Fig. 3, and Supplemental Table 1**).

74 Next, we performed a literature search of template plasmid integration at the target site in  
75 genome editing experiments, to determine the prevalence of this class of unintended  
76 alterations. We found that while there are reports, the template plasmid integration is often  
77 not a major finding, and thus we suspect that the integration errors are under reported or

78 overlooked. Template plasmid integration events are known to occur with zinc finger  
79 nucleases (ZFNs) at both the target and off-target sites<sup>10, 11</sup>. Using HEK-293 cells, Olsen, *et*  
80 *al.*<sup>10</sup> showed that transfection of plasmid alone resulted in plasmid integration at a rate of  
81  $28 \times 10^{-5}$  per cell; the addition of one ZFN increased the frequency to  $55 \times 10^{-5}$  per cell and  
82 two ZFNs further increased the frequency to  $99 \times 10^{-5}$  per cell. Work by Dickinson *et al.*<sup>12</sup>,  
83 using CRISPR/Cas9 with a template plasmid in *C. elegans*, reported the integration of a  
84 second copy of the template at the target site. Additional publications using CRISPR/Cas9  
85 with double stranded DNA (dsDNA) repair templates showed that the dsDNA templates can  
86 form multimers that integrate into the target site in fish<sup>13</sup> and mice<sup>14</sup>.

87 Our discovery highlights a potential blind spot in standard genome editing screening  
88 methods. In light of our finding, we propose modifications to current screening methods to  
89 enable detection of plasmid integration and integration of multiple template copies. The  
90 alignment of sequencing data should include both the reference genome and plasmid  
91 sequences. PCR genotyping should incorporate plasmid-specific primers. Methods to detect  
92 increased copies of the template and unintended integration of the template plasmid include  
93 long-range PCR conditions, quantitative PCR (e.g., digital droplet PCR), Southern blot, and  
94 long-read sequencing (e.g., Nanopore, PacBio). The application of suitable screening  
95 methods will provide a more precise measure of the prevalence of template plasmid  
96 integration events and drive improvements to genome editing, to the benefit of the field.

97 **Figures**



98

99 **Fig. 1: Template plasmid integration at the target site of genome-edited calves.**

100 Genomic structure of the template plasmid (**a**), unedited parental cell lines (**b**), and the  
101 genome-edited calves (**c**). (**a**) The repair template, containing the  $P_c$  sequence and flanking  
102 homology arms, is inserted in the pCR2.1 plasmid in an antisense orientation at the TOPO  
103 cloning site. (**b**) The unedited parental cell lines are homozygous for *HORNED*. (**a**) The  
104 genome-edited calves are heterozygous: one chromosome contains the intended edit ( $P_c$ ),  
105 while the other chromosome harbors template plasmid integration, in addition to the  
106 intended edit.

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