

1 The knockout of the HMG domain of the porcine SRY gene causes 2 sex reversal in gene-edited pigs

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43 **1 Abstract**

44 The sex-determining region on the Y chromosome (SRY) is thought to be the central genetic
45 element of male sex development. Mutations within the SRY gene are associated with a
46 male-to-female sex reversal syndrome in humans and other mammalian species such as
47 mice and rabbits. However, the underlying mechanisms are largely unknown. To understand
48 the biological function of the SRY gene, a site-directed mutational analysis is required to
49 investigate associated phenotypic changes at the molecular, cellular and morphological
50 level. In our study, we successfully generated a knockout of the porcine SRY gene by
51 microinjection of two clustered regularly interspaced short palindromic repeats (CRISPR) –
52 associated protein - 9 nuclease (Cas9) ribonucleoprotein (RNP) complexes targeting the
53 centrally located “high mobility group” (HMG) domain of the SRY gene. Mutations within this
54 region resulted in the development of complete external and internal female genitalia in
55 genetically male pigs. The internal female genitalia including uteri, ovaries, and oviducts,
56 revealed substantial size differences in 9-months old SRY-knockout pigs compared to age-
57 matched female wild type controls. In contrast, a deletion within the 5’ flanking region of the
58 HMG domain was not associated with sex reversal. Results of this study demonstrates for
59 the first time the central role of the HMG domain of the SRY gene in male sex determination
60 in pigs. Moreover, quantitative analysis by digital PCR revealed evidence for a duplication of
61 the porcine SRY locus. Our results pave the way towards the generation of boars exclusively
62 producing phenotypically female offspring to avoid surgical castration without anesthesia in
63 piglets. Moreover, the study establishes a large animal model that is much more similar to
64 humans in regard of physiology and anatomy and pivotal for longitudinal studies.

65

66 **2 Introduction**

67 In mammals, the male and female sex are determined by the presence or absence of the Y
68 chromosome (1). The sex-determining region on the Y chromosome (SRY) is located on the
69 short arm of the Y chromosome and is presumed to be critical for sex determination during
70 embryogenesis (2, 3). In pigs, the SRY gene consists of a single exon, with an open reading
71 frame of 624 bp representing 206 amino acids and encodes for the testis-determining
72 transcription factor (TDF). It is expressed in the male genital ridge at the time of sex
73 determination (4). The porcine SRY gene is first expressed on day 21 post coitum (p.c.) with
74 highest expression levels between day 21 and 23 p.c. Shortly after onset of SRY expression,
75 testis formation can be histologically determined between day 24 to 27 p.c. (5, 6).
76 Accordingly, the SRY gene is assumed to serve as the master regulator causing the formation
77 of primary precursor cells of tubuli seminiferi leading to the development of testicles from
78 undifferentiated gonads (7). However, it is still unknown whether the SRY gene is the only
79 sex-determining gene on the Y chromosome or if other genes such as SOX9 (8-10) and SOX3
80 (11) are involved as well.

81 In previous studies in mice (12) and rabbits (13), the SRY gene was knocked out using
82 different target regions. Both, the knockout of 92 % of the murine SRY gene by TALENs, or
83 the CRISPR/Cas-mediated knockout of the Sp1-DNA-binding sites of the rabbit SRY gene
84 caused sex reversal. Nevertheless, sequence divergence of the SRY gene between
85 mammalian species limited its direct structural and functional comparison and the
86 investigation of mammalian sex determination. So far, analysis of the SRY gene has almost
87 exclusively been done in small animals, mostly mice, and knowledge about the SRY gene in
88 large animal species, especially the porcine SRY gene is scarce.

89 The goal of the present study was to characterize the porcine SRY gene and its HMG domain
90 in male sex determination by knocking out different target sites of the porcine SRY gene via

91 intracytoplasmic microinjection of two CRISPR/Cas9 RNPs or cell transfection followed by
92 somatic cell nuclear transfer (SCNT) (Fig. 1). The generation of SRY-knockout pigs give
93 insights into the biological function of the SRY gene in a large animal species. While, the
94 murine SRY gene shows only 75% similarity to the human SRY gene, the porcine and human
95 SRY genes are closely related (~85 % amino acid homology) and show similar expression
96 profiles (5, 14). Therefore, a knockout of the highly conserved HMG domain in the porcine
97 model may pave the way for a suitable large animal model for the human male-to-female
98 sex reversal syndrome.

100 4 Results

101
102 **Production of SRY-knockout pigs.** To generate SRY-knockout (SRY-KO) pigs, a deletion of
103 approx. 300bp, encompassing the HMG domain in the porcine SRY gene was introduced (Fig.
104 2a). Thirty-one and thirty-two embryos derived from intracytoplasmic microinjection of
105 gRNA SRY_1 and SRY_3 into IVF-produced zygotes were surgically transferred into three
106 synchronized sows. Two recipients went to term and delivered twelve healthy piglets with
107 female phenotypes (Tab. 1/Fig. 3). Three of the piglets showed a deletion of approx. 300 bp
108 within the HMG domain of the SRY gene (Fig 4). Sequencing of the target region revealed
109 frameshift mutations of – 266 bp in piglet 715/2, and – 292 bp in piglet 715/7. Two different
110 genetic modifications, including a deletion of 298 bp and an indel formation with a deletion
111 of 298 bp and an insertion of 1 bp were detected in piglet 714/1 (Fig. 5).³⁴ Furthermore,
112 analysis of six Y chromosome specific genes (KDM6A, TXLINGY, DDX3Y, CUL4BY, UBA1Y and
113 UTY) demonstrated a male genotype and successful sex reversal in these piglets (Fig. S1, Tab.
114 S1). To ultimately confirm the male genotype of these piglets (715/2, 715/7 and 714/1), cells
115 from ear tissue were karyotyped detecting the Y chromosome in all three piglets (Fig. 6, Fig.
116 S2). No chromosomal abnormalities were observed in the sex-reversed pigs 715/2 and
117 715/7, while piglet 714/1 revealed an inversion of chromosome 7 (Fig. S2). The origin of this
118 clonal cytogenetic aberration remains unclear. It may not necessarily be related to the
119 CRISPR/Cas system because no off-target event was found on chromosome 7. In total, 34
120 potential off-target sites within the porcine genome were identified
121 (<http://crispor.tefor.net/>). We designed primers for the top ten off-target sites for each
122 gRNA (Tab. S2/S3). In one off-target site for gRNA_SRY1 and three off-target sites for
123 gRNA_SRY3 PCR amplification following Sanger sequencing was not possible. Overall, no off-
124 target events were observed (Fig. S3-S5). All SRY-KO pigs developed normally without any
125 health impairment (Fig. S6, Tab. S4).

126 In a second approach, one piglet generated via intracytoplasmic microinjection of gRNAs
127 SRY_1 and SRY_2 targeting the 5' flanking region of the HMG domain (Fig. 2b) was born with
128 an in-frame mutation of – 72 bp on the SRY locus, which did not lead to sex reversal (Fig. S7).

129
130 **External and internal genitalia of the SRY-KO pigs.** We investigated the external and internal
131 genitalia of the SRY-KO pigs. Age-matched wild type (WT) females from conventional
132 artificial insemination and female littermates of the SRY-KO pigs produced by microinjection
133 served as controls. At the age of 34 days, the external genitalia of the SRY-KO piglets were
134 similar to the external genitalia of female littermates and WT controls. To further investigate
135 the internal genitalia, the ovaries, oviducts and uteri of the 34 days old SRY-KO piglets and
136 female controls were prepared. The SRY-KO piglets had complete female internal genitalia,
137 including ovaries, oviducts and uteri that were similar to that of age matched WT females

138 (Fig. S8). Moreover, histological analysis of the inner structure of the ovaries revealed no
139 alteration in these young piglets (Fig. S9).
140 However, substantial size differences of the female genitalia were obvious in 9-months old
141 SRY-KO pigs compared to age-matched wild type controls (Fig. 7), with gene-edited animals
142 showing a substantially smaller genital tract. The SRY-KO pigs were not observed in heat,
143 even after three consecutive treatments of 1,000 IU PMSG (Pregmagon®, IDT Biologika)
144 followed 72 hours later by an intramuscular injection of 500 to 1,000 IU hCG (Ovogest®300,
145 MSD Germany) to induce estrus. Histological analysis of the ovaries of 9-months old SRY-KO
146 pigs revealed a high amount of loose connective tissue (Fig. S10). These results provided
147 evidence that the SRY-KO caused sex reversal in genetically male pigs with the development
148 of female external and internal genitalia, which lends further support to the central role of
149 the SRY gene in male sex determination during porcine embryogenesis. Re-cloning of piglet
150 715/2 led to seven sex-reversed piglets and demonstrated unequivocally that the elaborated
151 strategy described in this study can be used to successfully produce sex-reversed pigs (Fig.
152 S11 to S14).

153

154 **Duplication of the porcine SRY gene.** Investigation of the HMG domain of the porcine SRY
155 gene revealed one pig derived from intracytoplasmic microinjection (714/1) that displayed
156 two different genetic modifications within the SRY locus (Fig. 5). Whether these
157 modifications originated from mosaicism caused by the microinjection or a duplication of the
158 SRY locus was further analyzed. Analysis of different organ samples (liver, heart, colon,
159 kidney, spleen and lung) of piglet 714/1 revealed the same two genetic modifications in all
160 samples, arguing against mosaicism but for consideration of a duplicated SRY locus. Whole
161 genome sequencing using the USB-connected, portable Nanopore sequencer MinION
162 (Oxford Nanopore Technologies) and digitalPCR (QuantStudio®3D, ThermoFisher Scientific)
163 was performed to check for a possible SRY gene duplication. Due to the high number of
164 repetitive genes on the Y chromosome, Nanopore technology was used to sequence large
165 DNA fragments. However, only one contig similar to the SRY sequence could be found with
166 the assembled reads of the Nanopore Sequencing data. Even in a broad range, the sequence
167 in the flanking regions of the SRY locus was nearly identical hampering the analysis of the
168 duplicated SRY gene when using only the assembled Nanopore Sequencing data.

169 To further verify the duplication of the SRY gene, genomic DNA was analyzed by digital PCR.
170 Three targets, including the monoallelic SRY and KDM6A genes on the Y chromosome, and
171 the biallelic GGTA (galactosyltransferase) gene on chromosome 1 were selected for direct
172 comparison of their copy numbers. The copy numbers of GGTA1 were set to two (biallelic),
173 whereas the KDM6A and SRY genes were quantified in relation to the GGTA1 gene. In a first
174 trial, comparison of the copy numbers of the KDM6A and GGTA1 genes revealed a 2-fold
175 lower copy number of the monoallelic KDM6A compared to the biallelic GGTA1 in a male
176 wild type control (WT 7214 F2), as expected. In contrast, the normally monoallelic SRY gene
177 exhibited a similar calculated copy number as the biallelic GGTA1 gene (Fig. 8), indicating a
178 duplication of the SRY gene.

179 A second approach confirmed these findings by comparison of the copy numbers of the SRY
180 and GGTA1 gene in pigs with a complete SRY-KO (714/1, 715/2, 715/7), an incomplete SRY-
181 KO (713/1) and wild type controls (Fig. 9). As expected, no signal for SRY was detected in pigs
182 with a complete SRY-KO. For the incomplete SRY-KO, a piglet derived from intracytoplasmic
183 microinjection of plasmids SRY_1 and SRY_3 was used that showed two genetic
184 modifications, i.e. a 3 bp and a 298 bp deletion, within the SRY locus (Fig. S15/S16). Analysis
185 of several organ samples (liver, heart, colon, spleen, kidney, epididymis, testis and lung)

186 revealed the same genetic modification in all organs, indicating that mosaicism was highly
187 unlikely (Fig. S17). In this piglet, dPCR showed a 50 % reduced copy number of the SRY gene
188 compared to the GGTA1 gene. The SRY probe bound to the SRY locus with the smaller
189 deletion of 3 bp that did not interfere with the SRY assay and thereby indicated a duplication
190 of the SRY locus. As mentioned above, a similar copy number of the monoallelic SRY gene
191 compared to the biallelic GGTA1 gene was detected in wild type controls (Fig. 9).
192 To further exclude that these findings originated from mosaicism, two healthy piglets were
193 produced via SCNT using porcine fibroblasts edited with gRNAs SRY_1 and SRY_2 targeting
194 the 5' flanking region of the HMG domain of the SRY gene (Fig. 2b, Fig. S18) as donor cells.
195 Sequencing of the target site revealed two deletions of 72 bp and 73 bp in both piglets
196 (704/1 and 2) (Tab. S5, Fig. S19/S20). These results finally proved the presumptive
197 duplication of the porcine SRY locus.

198

199 **5 Discussion**

200

201 The sex-determining region on the Y chromosome (SRY) is critically involved in mammalian
202 male sex development (5). However, the molecular function and the role that the SRY gene
203 plays as the main switch for male sex development in all mammals are yet to be explored.
204 Previous studies in mice (12) and rabbits (13) investigated the potential role of the SRY gene
205 for sex development. The murine SRY gene was knocked out by introducing two base pairs
206 into the 5' part of the ORF (open reading frame) of the SRY gene causing a frameshift. In one
207 genetically male offspring this mutation led to a female phenotype (12). In rabbits, a
208 disruption of the Sp1-binding site in the 5' flanking region of the SRY gene also resulted in
209 sex reversal (13). In contrast, an in-frame mutation upstream of the HMG domain of the
210 porcine SRY gene described in this study did not result in the generation of genetically male
211 offspring with a female phenotype. Detection of two genetic modifications in several pigs
212 provided evidence of a presumed duplication of the porcine SRY locus. Skinner et al.
213 described the porcine SRY gene in a two palindromic head-to-head copy manner (15), as in
214 rabbits (16). Quantitative analysis by digital PCR (QuantStudio®3D, ThermoFisher Scientific)
215 revealed duplication of the SRY locus by detection of a similar copy number of the
216 monoallelic SRY and the biallelic GGTA1 genes. Moreover, a reduction of the copy number of
217 the SRY gene from wild type control to a complete SRY-KO pig displayed the presumed
218 duplication in pigs generated via intracytoplasmic microinjection. Ultimately, the generation
219 of pigs via SCNT that carried two different deletions within the SRY gene confirmed the
220 presence of the SRY duplication, because cloning technique avoids any mosaicism. Nanopore
221 sequencing indicated a high similarity of the two SRY loci impairing the differentiation of
222 both loci. The alignment of the assembled reads to the reference sequence resulted in loss
223 of information that might be crucial for differentiation of both SRY loci. An evaluation of the
224 raw data, a de-novo assembly or an enhancement of the Nanopore Sequencing data with
225 Illumina MiSeq data could overcome these limitations (17-19). It is still unknown, whether
226 both copies of the porcine SRY gene are active and required for male sex development and if
227 there is the need to reach a certain threshold expression level from the SRY locus to induce
228 the development of a male gender as previously described in mice (20-22). To address these
229 questions, the identification of potential single nucleotide polymorphisms (SNPs) to
230 differentiate between the two SRY loci is desirable.

231 We report here for the first time the successful knockout of the HMG domain of the porcine
232 SRY gene by intracytoplasmic microinjection of two CRISPR/Cas9 RNP complexes resulting in
233 genetically male pigs with a female phenotype. The CRISPR/Cas9 system has emerged as the

234 genome editing technology of choice for many applications due to its ease of use, cost-
235 effectivity and high specificity to introduce mutations at the targeted loci (23, 24).
236 Nevertheless, off-target cleavages at undesired genomic sites may occur. It is necessary to
237 further increase the specificity of the CRISPR/Cas system regarding the gRNA design (25), by
238 involving CRISPR nickase proteins (26), using anti-CRISPR proteins (27), employing
239 ribonucleoproteins (RNPs) (28, 29) or designing “self-restricted” CRISPR/Cas systems (30).
240 CRISPR/Cas9 RNP components persist only temporarily in cells thereby limiting guideRNA
241 and Cas9 expression to a short time window. The use of CRISPR/Cas9 RNPs enables efficient
242 genome editing while significantly reducing possible off-target events and mosaicism
243 formation. Random integration of DNA segments into the host genome as with DNA
244 plasmids is avoided by using RNPs (28, 29, 31). However, no off-target events were found at
245 possible sites using PCR-based analysis and Sanger sequencing in the SRY-KO pigs generated
246 via intracytoplasmic microinjection of CRISPR/Cas RNPs. However, only with whole-genome
247 sequencing using accurate and sensitive off-target profiling techniques such as GUIDE-Seq
248 and CIRCLE-Seq the occurrence of unexpected mutations could be excluded completely (25,
249 32, 33).

250 In the present study, healthy SRY-KO pigs showing normal development and growth rates
251 were born (Fig. S6, Tab. S4). Moreover, the knockout of the HMG domain resulted in piglets
252 with a female phenotype, including female external and internal genitalia. A previous study
253 reported that rabbits with a knockout in the Sp1-binding sites of the SRY gene showed a
254 dramatically reduced number of follicles in their ovaries (13). Although a normal copulatory
255 behavior was observed, no pregnancy was established by mating of the genetically modified
256 rabbits to wild type male rabbits. Transfer of blastocysts from wild type female rabbits into
257 pseudo-pregnant SRY-KO rabbits resulted in a successful pregnancy with the birth of twelve
258 pups. It was assumed that the abnormal development and reduced number of follicles were
259 responsible for the decreased fertility in the sex-reversed rabbits. In our study, substantial
260 size differences in all female genitalia of 9-months old SRY-KO pigs compared to the age-
261 matched wild type controls demonstrated markedly retarded development of female
262 genitalia. It has to be clarified, whether Y chromosome induced gene and hormone
263 expression hampered the development of female genitalia in those SRY-KO pigs. One
264 example of the influence on female sex development from disturbed hormone profiles
265 (androstenedione and müllerian inhibition substance) in females is the bovine freemartin
266 syndrome which leads to the masculinization of the female genitalia (34). Moreover, the
267 absence of the second X chromosome in the SRY-KO pigs might have an impact on female
268 sex development (35). The inactivation of one copy of the X chromosome is essential for
269 undisturbed female development, nevertheless, several genes (mainly located on the short
270 arm of the X chromosome) usually escape X chromosome inactivation (36). Further studies
271 are necessary to investigate the gene expression and hormone levels in these SRY-KO pigs.
272 An XO phenotype lacking the Y chromosome would be a promising animal model to
273 investigate the influence of Y chromosomal gene expression and to clarify the importance of
274 the second X chromosome in female sex development. It was previously shown, that
275 CRISPR/Cas-mediated elimination of the murine Y chromosome is possible by targeting a
276 cluster of genes along the Y chromosome (37). Moreover, in human embryonic stem cells it
277 was shown that the CRISPR/Cas3 system has the potential to induce long-range
278 chromosomal deletions (38). Both methods can be utilized to generate a porcine XO
279 phenotype.

280

281 The results of this study further clarified the critical role of the porcine SRY gene in male sex
282 determination. The pre-determination of sex using CRISPR/Cas9 targeting the porcine SRY
283 gene could be of great benefit for animal welfare elimination the need for castration of male
284 offspring to avoid boar taint by delivery of phenotypically female piglets. Currently, most
285 piglets are surgically castrated without anesthesia shortly after birth, which raised animal
286 welfare concerns and resulted in a ban of this practice within the EU. It was recently
287 reported that knockout of the KISSR gene by TALEN-mediated mutagenesis resulted in the
288 generation of boars that remained in the pre-pubertal stage lacking boar taint (39).
289 However, to use these animals for breeding purposes, they have to be hormonally treated,
290 which in turn might result in reduced consumer acceptance. Our results could pave the way
291 for the production of boars that produce only female offspring by integrating a CRISPR/Cas9
292 vector targeting the HMG domain of the SRY gene into the porcine genome. The transgenic
293 founder would produce feminized males (XY^{SRY^-}) and normal females. Alternatively, the
294 CRISPR/Cas vector could target multiple genes on the Y chromosome during
295 spermatogenesis to prevent development of Y-chromosomal sperm. Thereby, only female
296 offspring would be generated. In both approaches, use of a self-excising vector should result
297 in the generation of non-transgenic offspring. It remains to be determined whether products
298 from genome-edited animals will find market acceptance in light of a critical public debate
299 on genome engineering in many countries. Nevertheless, the above-mentioned strategies
300 might improve welfare in pig farming and may lead to a more sustainable pork production.
301 In addition to its importance for animal welfare, the SRY-KO pigs could be useful for better
302 mechanistic insights into the human male-to-female sex reversal syndrome (Swyer
303 syndrome) (40). Overall, 15 to 20 % of humans exhibiting male-to-female sex reversal
304 syndrome carry mutations in or dysfunctions of the SRY gene. Most of the detected
305 variations in humans are located within the “high mobility group” (HMG) domain of the SRY
306 gene that is responsible for DNA binding (11, 41) and thought to act as the main functional
307 domain for SRY protein synthesis (42-44). The murine SRY gene shows only 75% similarity to
308 the human SRY gene. In contrast, the porcine and human SRY genes are closely related (~85
309 % amino acid homology) and show similar expression profiles (5, 14). Taken this in account,
310 the high similarity of the HMG domain and the high degree of physiological, genetic and
311 anatomical similarity of the pig to humans renders the pig as a promising large animal model
312 to gain insight into human sex determination and the interaction of sex chromosome related
313 gene expression profiles (5, 45).

314

315 **6 Materials and Methods**

316 **Animal Welfare.** Animals were maintained and handled according to the German guidelines
317 for animal welfare and the genetically modified organisms (GMO) act. The animal
318 experiments were approved by an external animal welfare committee (Niedersaechsisches
319 Landesamt fuer Verbraucherschutz und Lebensmittelsicherheit, LAVES, AZ: 33.9-42502-04-
320 17/2541), which included ethical approval of the experiments.

321 **Transfection of gRNAs.** The CRISPR/Cas9 system was employed to induce defined deletions
322 within the SRY gene (Ensembl transcript: ENSSSCG00000037443). Guide RNAs (gRNAs)
323 targeting either the 5' flanking region of the HMG domain of the SRY gene (SRY_1 and
324 SRY_2) or encompassing the HMG box (SRY_1 and SRY_3) were designed using the web-
325 based design tool *CRISPOR* (<http://crispor.tefor.net/>) (Fig.2). Target sequences were further
326 analyzed via BLAST to reduce the probability for off-target events. The gRNA oligos with a
327 BbsI overhang were cloned into the linearized CRISPR/Cas9 vector pX330 (addgene, #42230).

328 Afterwards, two CRISPR/Cas9 plasmids were co-transfected (with a final concentration of 5
329 $\mu\text{g}/\mu\text{l}$) into male porcine fibroblasts by electroporation (NeonTM Transfection System,
330 ThermoFisher Scientific) to test the efficacy of the plasmids to induce double-strand breaks
331 at the targeted locus. Electroporation conditions were as follows: 1350 V, 20 mm, and two
332 pulses. After lysis of transfected cells, the cell lysate was analyzed using SRY specific primer
333 (SRY-F: 5'-TGAAAGCGGACGATTACAGC and SRY-R: 5'-GGCTTTCTGTTCTGAGCAC-3'). The
334 purified PCR product (10 $\text{ng}/\mu\text{l}$) (Invisorb[®] Fragment CleanUp – Startec) was Sanger
335 sequenced to detect mutations at the target site.

336 **In-Vitro-Fertilization and In-Vitro-Maturation.** In-vitro-maturation of porcine oocytes was
337 performed as previously described (46). Frozen boar semen from a fertile landrace boar was
338 thawed for 30 sec. in a water bath (37 °C). The motility of sperm was analyzed using
339 microscopy (Olympus, BH-2). After washing with Androhep[®] Plus (Minitube) and
340 centrifugation for 6 minutes at 600 g, approx. 75 to 100 sperm per oocyte (depending on
341 semen capacity) were used for fertilization (no sexed sperm were utilized for fertilization).
342 After four hours of co-incubation, the fertilized oocytes were cultured in porcine-zygote-
343 medium (PZM-3 medium).

344 **Somatic cell nuclear transfer.** SCNT was performed as previously described (47). Fetal
345 fibroblasts transfected with gRNA SRY_1 and SRY_2 targeting the flanking region of the HMG
346 domain of the SRY gene were used as donor cells. Eighty-two and eighty-six one- to two-cell
347 embryos were surgically transferred into two hormonally synchronized German Landrace
348 gilts (7 to 9-months old). Estrus was synchronized by application of 20 mg/day/gilt
349 Altrenogest (Regumate[®] 4mg/ml, MSD Germany) for 12 days, followed by an injection of
350 1,500 IU PMSG (pregnant mare serum gonadotropin, Pregmagon[®], IDT Biologika) on day 13
351 and introduction of ovulation by intramuscular injection of 500 IU hCG (human
352 choriongonadotropin, Ovogest[®]300, MSD Germany) 78 h after PMSG administration.

353 **Preparation of RNP complexes for microinjection.** The Alt-R CRISPR/Cas9 system (IDT)
354 consists of two CRISPR RNA components (crRNA and tracrRNA). The crRNA was individually
355 designed to target the HMG domain of the SRY gene (SRY_3: 5' – AAATACCGACCTCGTCGCAA
356 – 3'). To generate an active gRNA, both components (crRNA and tracrRNA) were annealed (95
357 °C for 5 min and then ramped down to 25 °C at 5 °C/min) in a ratio of 1 : 1 to reach a final
358 concentration of 1 $\mu\text{g}/\mu\text{l}$. Afterwards, the gRNA complex was mixed with Alt-R S.p. Cas9
359 nuclease 3NLS and incubated for 10 minutes at room temperature to form an active RNP
360 complex with a final concentration of 20 $\text{ng}/\mu\text{l}$. The second RNP complex was prepared using
361 the individually designed synthetic single-guide RNA (SRY_1: 5' – ATTGTCCGTCGGAATAGTG
362 – 3') from Synthego. The sgRNA was mixed with purified 2NLS-Cas9 nuclease using a ratio of
363 approximately 1 : 1.5 (0,84 μl sgRNA [25pmols] and 1.25 μl Cas9 protein [25 pmols]) and
364 incubated for 10 minutes at room temperature. After centrifugation at 10,000 rpm for 10
365 minutes and 4 °C, the supernatant was transferred into a new tube. Both RNP complexes
366 were mixed in a ratio of 1 (SRY_1) to 1.7 (SRY_3) and directly used for microinjection.

367 **Microinjection.** The RNPs targeting the SRY gene were intracytoplasmically co-injected
368 into IVF-produced zygotes obtained from slaughterhouse ovaries. Therefore, approx. 10 pl of
369 the RNP solution was injected with a pressure of 600 hPa into IVF-produced zygotes
370 (FemtoJet, Eppendorf). The injected zygotes were cultured in PZM-3 medium at 39 °C, 5 %
371 CO₂ and 5 % O₂. At day 5, when embryos reached the blastocyst stage, 31 or 32 embryos,
372 respectively, were surgically transferred into two recipients.

373 **Establishing cell cultures from SRY-KO piglets.** Porcine fibroblasts were isolated from ear
374 tissue of the piglets and cultured in Dulbecco's modified Eagle's medium (DMEM) with 2 %
375 penicillin/streptomycin, 1 % non-essential amino acids and sodium pyruvate and 30 % fetal
376 calf serum (FCS) (Gibco, 10270-106). When cells reached confluency, they were lysed with
377 EDTA/Trypsin and genomic DNA was analyzed by PCR and karyotyping.

378 **PCR-based genotyping.** Genomic DNA of the pigs was extracted from tail tips. Cells were
379 isolated from ear tissue. The DNA concentration was determined using the NanoDrop™
380 (Kikser-Biotech) system. For genotyping of the pigs, polymerase chain reaction (PCR) was
381 employed using specific primer (SRY-F: 5'-TGAAAGCGGACGATTACAGC-3' and SRY-R: 5'-
382 GGCTTTCTGTTCTGAGCAC-3') flanking a 498 bp segment of the SRY gene (Fig. 2). PCR
383 amplification was performed in a total volume of 50 µl : 20 ng DNA, 0.6 µM reverse and
384 forward primer, 1.5 mM MgCl₂, 0.2 mM dNTPs and 1.25 U *Taq* Polymerase. Cycling
385 conditions were as follows: 32 cycles with denaturation at 94°C for 30 sec, annealing at 59 or
386 60 °C for 45 sec, extension at 72°C for 30 sec and a final extension at 72°C for 5 minutes. The
387 standard conditions for gel electrophoresis were set up to 80 V, 400 mA and 60 min using a 1
388 % agarose gel. The PCR-product was purified (Invisorb®Fragment CleanUp-Kit, Startec) and
389 Sanger sequenced. To further analyze the genotype of the piglets Y chromosome specific
390 genes such as KDM6A, DDX3Y, CUL4BY, UTY, UBA1Y and TXLINGY were amplified
391 (Supplements Table 1).

392 **Karyotyping of the cells.** Karyotyping was accomplished on porcine fibroblasts isolated from
393 ear tissue. After treatment of cells for 30 minutes with colcemide (Invitrogen), cells were
394 trypsinized and metaphases were prepared according to standard procedures. Fluorescence
395 R-banding using chromomycin A3 and methyl green was performed as previously described
396 in detail (48). At least 15 metaphases were analyzed per offspring. Standard karyotype of the
397 pig includes 38 chromosomes. Karyotypes were described according to Gustavsson, 1988
398 (49) and the International System for Human Cytogenetic Nomenclature (ISCN).

399 **Histology.** Porcine ovarian tissues were fixed with 4 % paraformaldehyde for 6 to 8 hours
400 (smaller tissues of up to 5 x 10 mm) or overnight (tissues of up to 2 x 3 cm), incubated in 30
401 % sucrose for two hours and frozen at - 80 °C. Afterwards, the tissues were embed in
402 TissueTek® (Sakura, TTEK), cut in thin sections (15 µm) and stained with hematoxylin and
403 eosin (HE) following standard procedures (50). Analyzes of inner structure of ovaries were
404 done by microscopy (DMIL LED, Leica).

405 **Off-target analysis.** The top ten off-target effects were selected from the gRNA design tool
406 *CRISPOR* (<http://crispor.tefor.net/>). PCR primers used for amplifying the PCR product are
407 listed in Supplements Table 3 for SRY_1 and Supplements Table 4 for SRY_3. The PCR
408 product was purified (Invisorb®Fragment CleanUp-Kit, Startec, Germany) and analyzed via
409 Sanger sequencing.

410 **DigitalPCR.** Three assays including a probe and two primers (in a ratio of 2.5 probe to 9 nM
411 primer) targeting the SRY and KDM6A genes (FAM™-labeled) on the Y chromosome and
412 GGTA gene (HEX™-labeled) on chromosome 1 (as control) were designed (IDT) for digital
413 polymerase chain reaction (dPCR). The dPCR was performed in a total reaction volume of
414 14.5µl with the following components: 7.3 µl Master Mix (QuantStudio™3D Digital PCR
415 Master Mix v2, ThermoFisher Scientific), 0.7 µl HEX™ and VIC™ dye-labeled assays each, 1.4
416 µl diluted genomic DNA and 4.4 µl nuclease-free water. Standard dPCR thermal cycling
417 conditions were used with an annealing temperature of 60 °C in the QuantStudio™ 3D

418 Digital device (ThermoFisher Scientific). Copy numbers of the genes within each chip were
419 compared and analyzed via the QuantStudio™ 3D AnalysisSuite software
420 (<http://apps.lifetechnologies.com/quantstudio3d/>). The copy number of the GGTA1 gene
421 was set at 2 (biallelic), copy numbers of KDM6A and SRY genes were given in proportion to
422 the GGTA1 gene. All findings were verified in three replicates with variable DNA
423 concentration and different samples (51).

424 **Nanopore Sequencing.** Whole genome sequencing was performed by using the MinION
425 device of Oxford Nanopore Technologies to investigate the porcine SRY locus. DNA from a
426 male wild type blood sample (2 ml) was purified with the NucleoBand®HMW DNA Kit
427 (Macherey-Nagel). To eliminate fragments below 40 kb the Short Read Elimination Kit XL
428 (Circulomics) was utilized. Subsequently, 47 µl high molecular weight DNA (30 - 40 ng/µl)
429 was prepared with the Ligation Sequencing Kit (SQK-LSK109, Oxford Nanopore) and the
430 NEBNext® Companion Module for Oxford Nanopore Technologies® Ligation Sequencing
431 (BioLabs, E7180S) using the Nanopore Oxford standard protocol for ligation sequencing.

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433

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444 42230 ; <http://n2t.net/addgene:42230> ; RRID:Addgene_42230).

445

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575

576 **9 Figures and Tables**

577 **Figure 1** Schematic illustration of the experimental design to generate SRY-KO pigs (XY^{SRY⁻}) by either
578 intracytoplasmic microinjection of two CRISPR/Cas9 RNP complexes into IVF-produced zygotes or
579 somatic cell nuclear transfer (SCNT). Embryos were surgically transferred into hormonally
580 synchronized recipients, and the offspring were analyzed pheno- and genotypically.

581

582

583 **Figure 2 a** Location of two sgRNA target sites (yellow underlined) flanking the HMG-box (red box) of
584 the SRY gene. **b** Schematic illustration showing the guide RNAs (yellow underlined) targeting an
585 approx. 72 bp segment in the 5' flanking region of the HMG domain (red box) of the SRY gene. Primer
586 amplifying the SRY exon are indicated with green arrows.

587

588 **Figure 3** Twelve healthy piglets were born after cytoplasmic microinjection of two CRISPR/Cas9 RNP
589 complexes into IVF-produced zygotes and surgical embryo transfer. Three of the piglets showed
590 complete female external genitalia. The deletion of the SRY gene had no effect on growth rate
591 compared to wild type. All piglets developed normally.

592

593

594 **Figure 4** PCR-based detection of the mutated SRY gene in piglets (714/1 and 715/1-11) generated via
595 microinjection of CRISPR/Cas9 RNP complexes. Three piglets (715/2, 715/7 and 714/1, indicated with
596 white asterisk) showed deletions of approx. 300 bp within the SRY gene compared to a male wild
597 type control (WT 578 F7). The male WT control showed an expected band of ~500 bp. The female WT
598 control (WT 578 F4) is negative, as expected for the SRY gene.

599

600 **Figure 5** Sanger sequencing of the purified PCR product of the SRY-KO piglets (715/2, 715/7 and
601 714/1) showed genetic modifications within the SRY locus. Piglet 715/7 displayed a deletion of 292bp
602 and piglet 715/2 of 266bp. Piglet 714/1 showed two different mutations with a deletion of 298bp
603 and an indel formation of -298bp and +1bp.

604

605 **Figure 6** Karyotyping of cells from the SRY-KO piglet 715/2 confirmed the male genotype of this
606 piglet. The karyotypes of piglet 715/7 and 714/1 are shown in the Supplements Fig. 6.

607

608 **Figure 7** The uteri and ovaries of the 9-months old SRY-KO, XY pig (714/1) and the age-matched
609 WT,XX piglet (control from same litter). **a** Substantial size differences were displayed in the 9-months

610 old SRY-KO pig compared to the female wild type control. **b** The ovaries in the 9-month old SRY-KO,
611 XY pig were significant smaller than the ovaries of the WT, XX pig and showed no follicles.
612

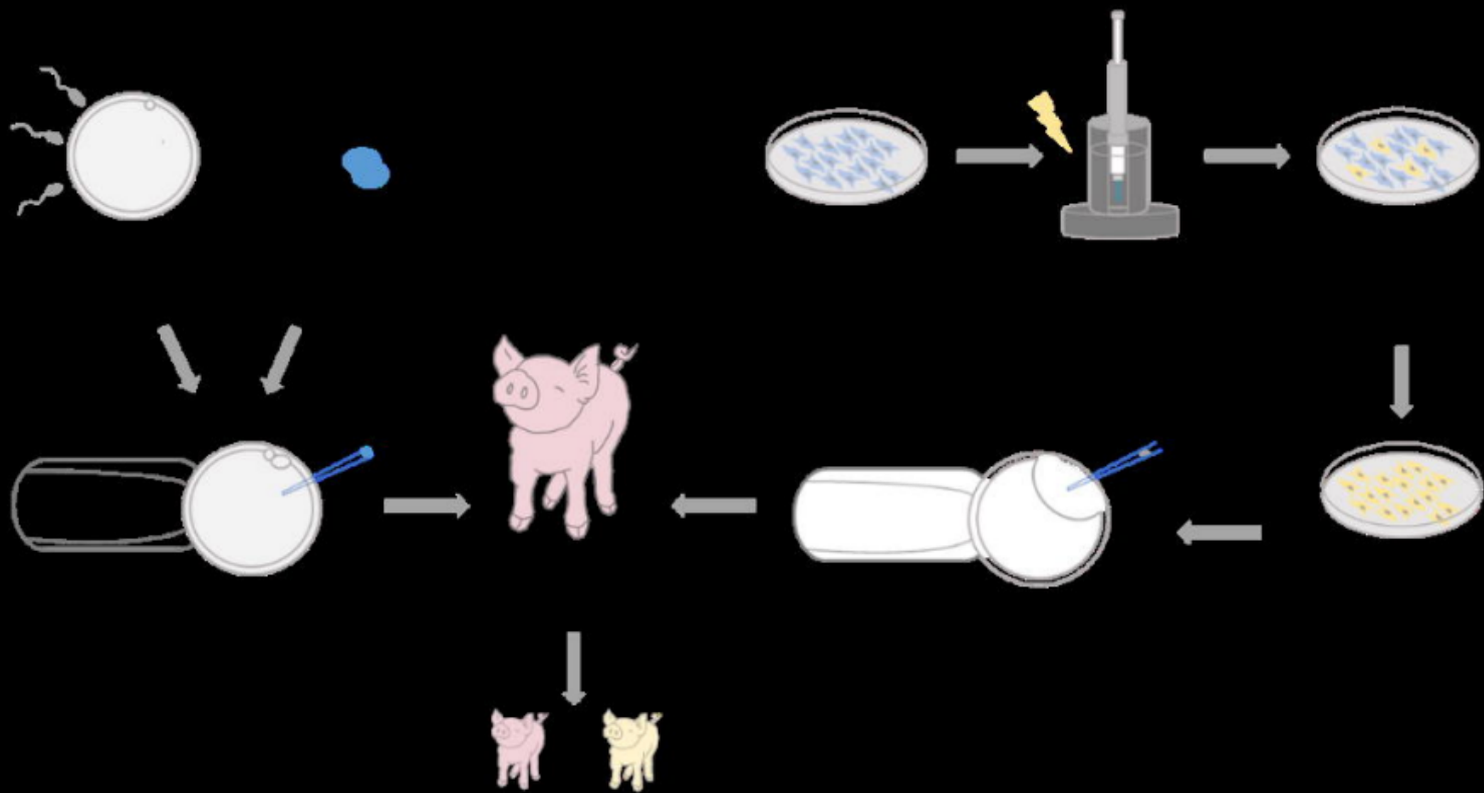
613 **Figure 8** Schematic diagram of the first trail of dPCR. The dPCR biplex assay revealed a two fold lower
614 copy number of the monoallelic KDM6A gene compared to the biallelic GGTA1 gene, as expected. A
615 similar copy number of the monoallelic SRY gene compared to the biallelic GGTA1 gene indicated a
616 duplication of the SRY locus.
617

618

619 **Figure 9** Schematic diagram of the second trail of dPCR. The dPCR biplex assay revealed a stepwise
620 reduction of the copy number of the SRY gene from wild type control to complete SRY-KO pig
621 compared to the GGTA1 gene. These results supported the assumption of a porcine SRY duplication.
622

623

624 **Table 1** Results of the embryo transfer of microinjected zygotes into recipients. Three of twelve
625 piglets showed a sex reversal with a female phenotype and a male genotype.



Exon SRY gene

HMG-box SRY gene

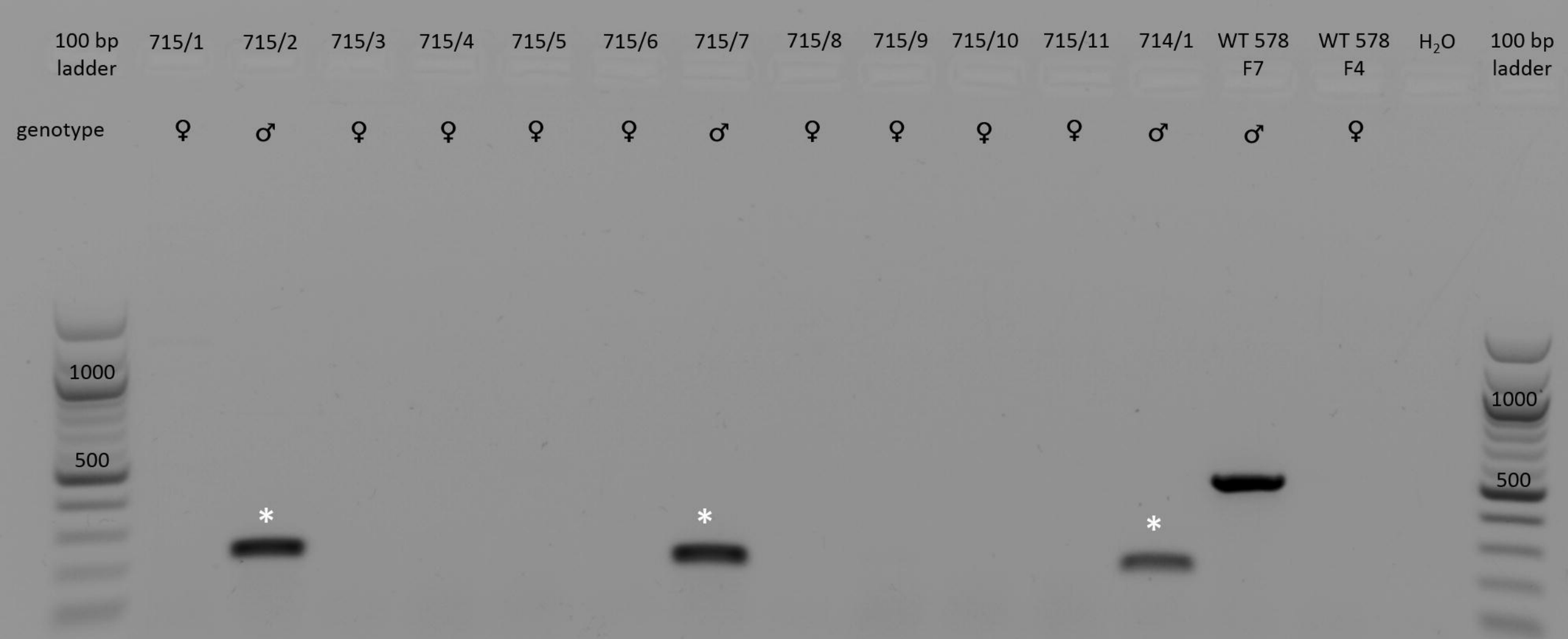
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CTCCTCACTATTTCCGACGGACAAT AAATACCGACCTCGTCGCAAGGG

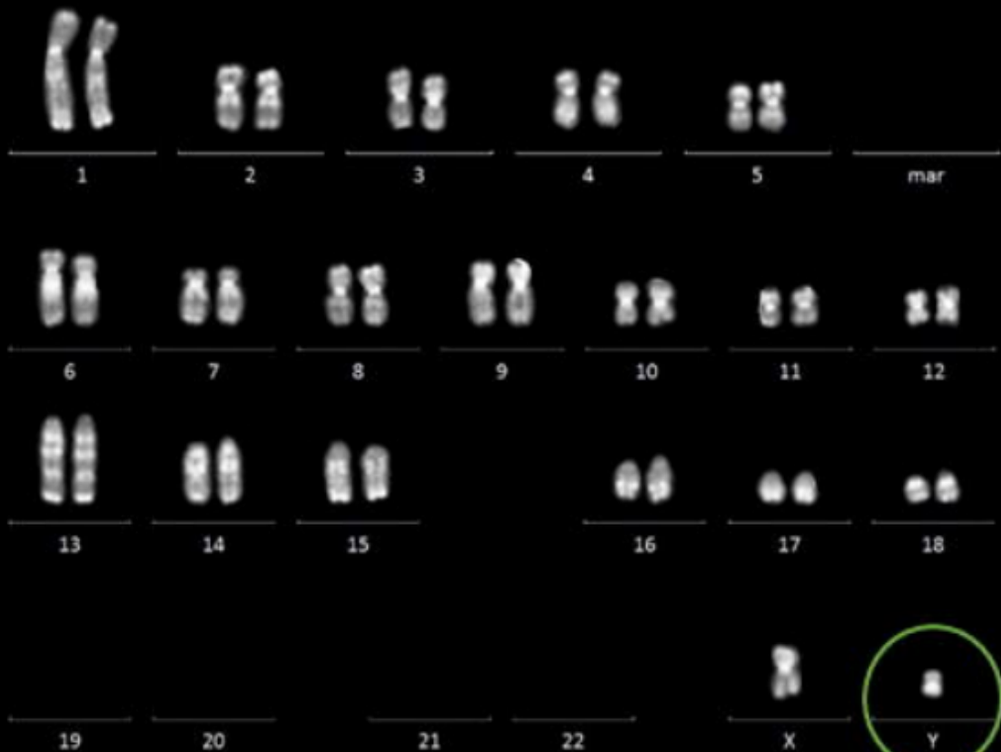
Exon SRY gene

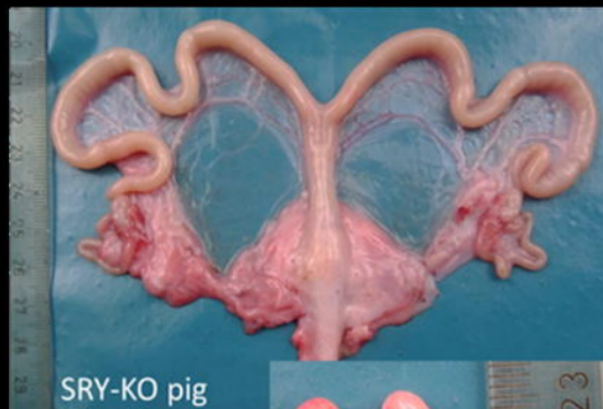
HMG-box SRY gene

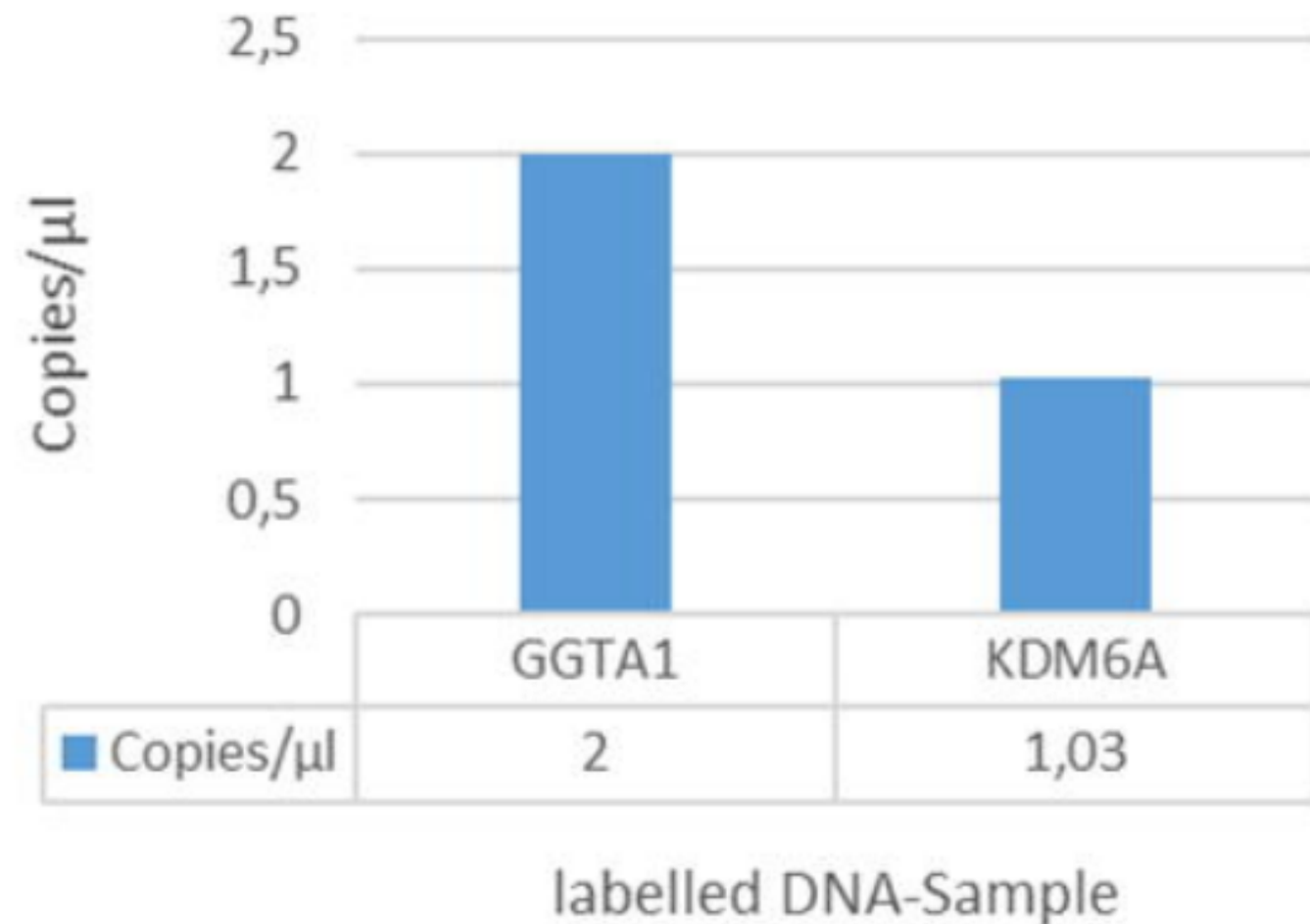
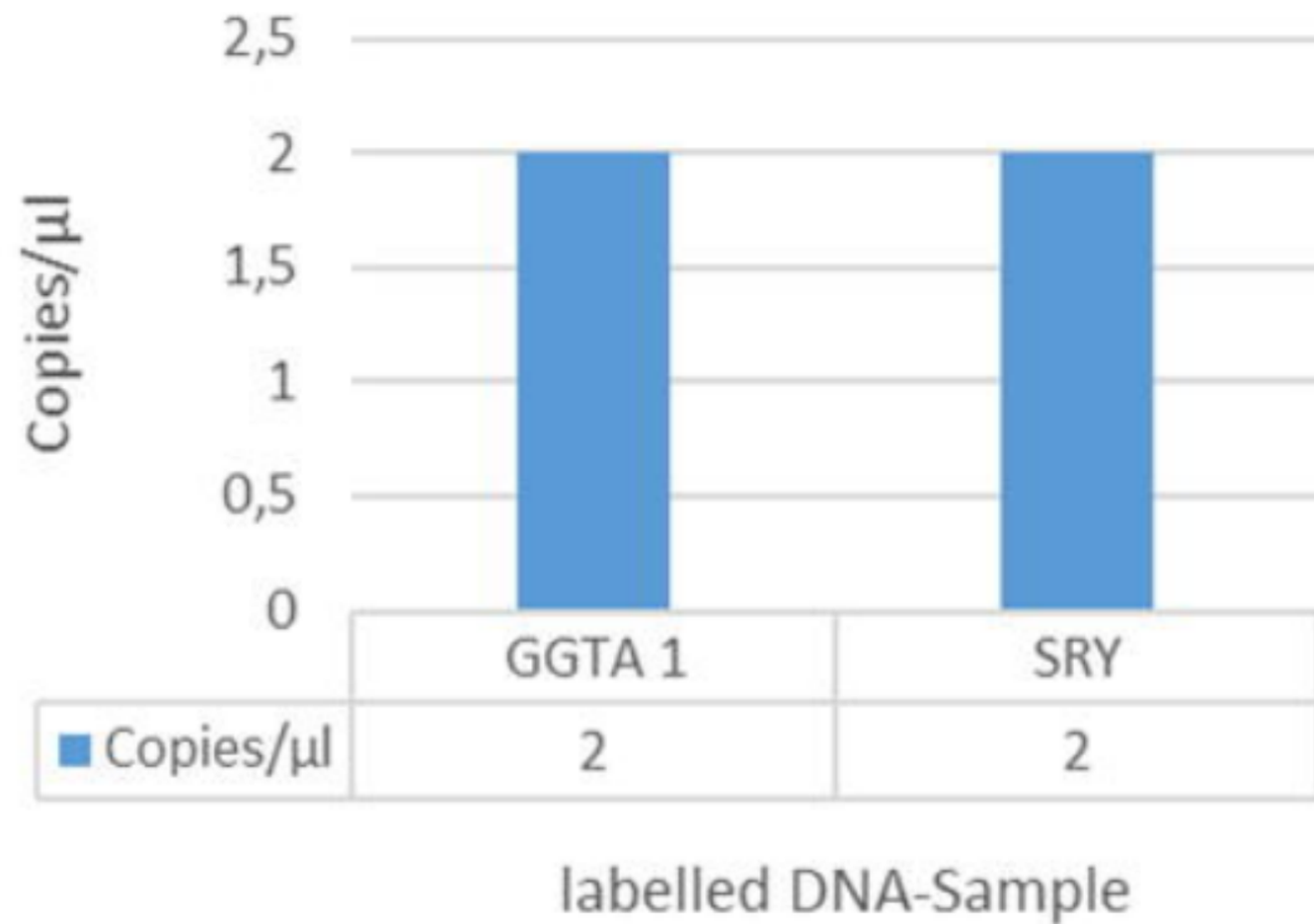
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CTCCTCACTATTTCCGACGGACAAT CCAGGATCGTGTCAAGCGACCCA

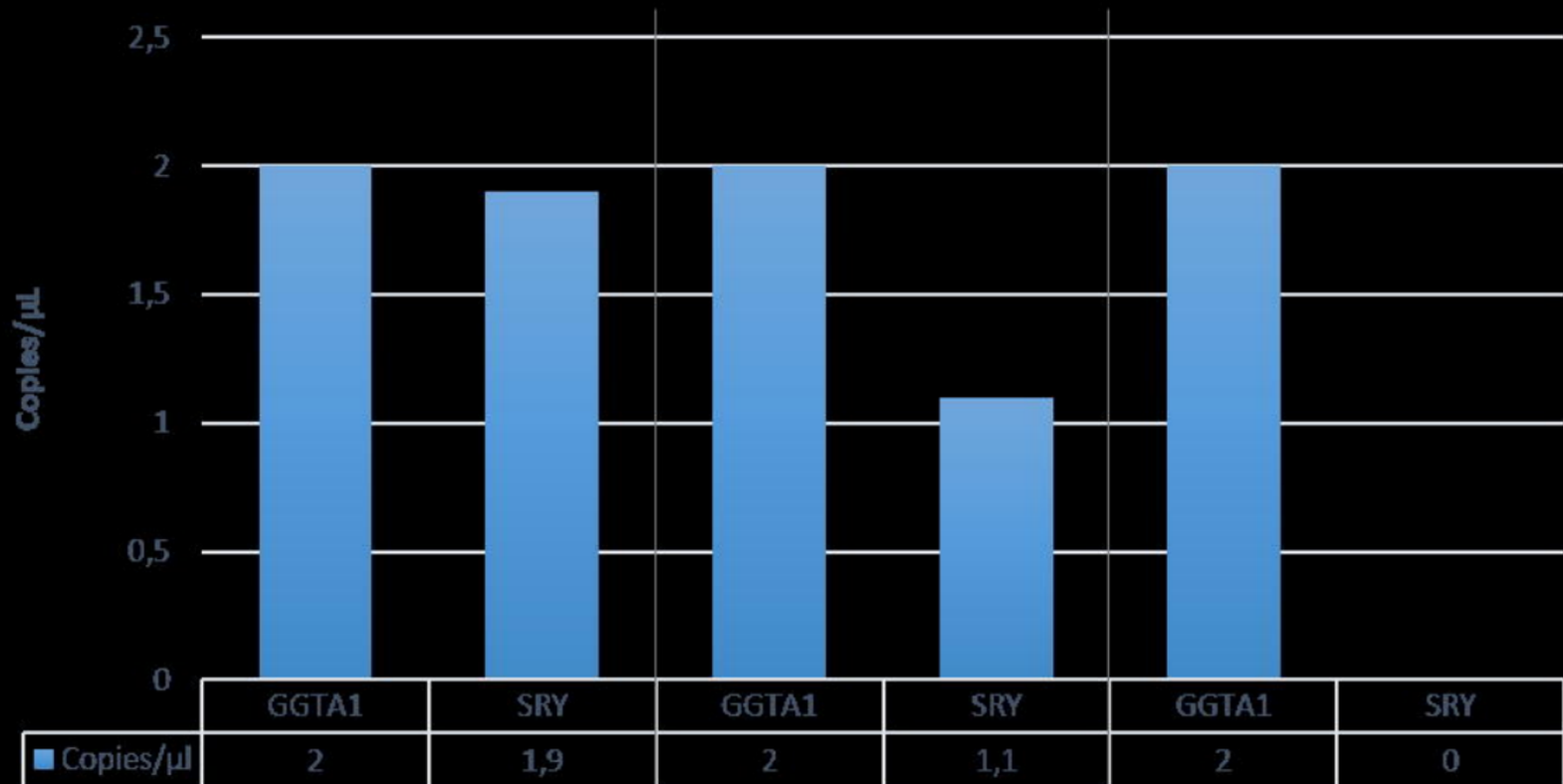












8018	32	-	-	-	-
715	31	+	11	2	2