1 Title

- 2 The Vertebrate Codex Gene Breaking Protein Trap Library For Genomic Discovery and Disease
- 3 Modeling Applications
- 5 Authors

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- SCE and KJC conceived research; NI, SCE, KJC for basic experimental design including detailed
- analyses of the collection; MS, CD, MU, SER, YD, RU, MM, JJE, XX, DB, SCE and KJC generated
- GBT collection, NI, KS, MS, LG, CD, MU, RU, YD, SAF, WL and KJC conducted phenotype
- screening of GBT mutant lines; NI, KS, MS, LG, CD, MU, RU, YD, WL and KJC conducted molecular
- biology analyses; GV and SB conducted next generation sequencing; NI, KS, SCE and KJC conducted
- 19 bioinformatics-based analyses; LAS, NI, KJC and SCE conducted comparative genomics analyses, NI,
- 20 KS, MS, LG, KJS and SCE wrote the manuscript; SCE, KJC, JJE, XX, MH, SAF, XYW, SB, XX and
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- 22 Keywords

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- 1 Gene-break transposon, zebrafish, disease model, human genetic disorders, gene ontology, Lightsheet
- 2 microscopy

Abstract

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The zebrafish is a powerful model to explore the molecular genetics and expression of the vertebrate genome. The gene break transposon (GBT) is a unique insertional mutagen that reports the expression of the tagged member of the proteome while generating Cre-revertible genetic alleles. This 1000+ locus collection represents novel codex expression data from the illuminated mRFP protein trap, with 36% and 87% of the cloned lines showcasing to our knowledge the first described expression of these genes at day 2 and day 4 of development, respectively. Analyses of 183 molecularly characterized loci indicate a rich mix of genes involved in diverse cellular processes from cell signaling to DNA repair. The mutagenicity of the GBT cassette is very high as assessed using both forward and reverse genetic approaches. Sampling over 150 lines for visible phenotypes after 5dpf shows a similar rate of discovery of embryonic phenotypes as ENU and retroviral mutagenesis. Furthermore, five cloned insertions were in loci with previously described phenotypes; embryos homozygous for each of the corresponding GBT alleles displayed strong loss of function phenotypes comparable to published mutants using other mutagenesis strategies (ryr1b, fras1, tnnt2a, edar and hmcn1). Using molecular assessment after positional cloning, to date nearly all alleles cause at least a 99+% knockdown of the tagged gene. Interestingly, over 35% of the cloned loci represent 68 mutants in zebrafish orthologs of human disease loci, including nervous, cardiovascular, endocrine, digestive, musculoskeletal, immune and integument systems. The GBT protein trapping system enabled the construction of a comprehensive protein codex including novel expression annotation, identifying new functional roles of the vertebrate genome and generating a diverse collection of potential models of human disease.

Introduction

With the generation of more than 100 sequenced vertebrate genomes (Meadows & Lindblad-Toh, 2017),
the current key question is how to determine the role(s) of uncharacterized gene products in specific
biological and pathological processes. For example, genes associated with human disease are being
discovered at a rapid rate. However, the biological functions underlying this linkage is often unclear
(Kettleborough et al., 2013). Model system science using loss of function approaches has been essential
to the annotation of the genome to date including the discovery of novel processes and the biological

mechanisms underlying disease (Stoeger, Gerlach, Morimoto, & Nunes Amaral, 2018).

Among vertebrates, *Danio rerio* (zebrafish) has emerged as an outstanding model organism amenable to both forward and reverse genetic approaches. In addition, the natural transparency of the zebrafish embryo and larvae enables the unprecedented ability to non-invasively collect a rich set of expression data for the proteome and in the context of an entire living vertebrate. We describe here a 1000+ collection of zebrafish lines made using the Protein Trap Gene- Breaking Transposon (GBT;(Clark, Balciunas, et al., 2011)to develop such a codex for the comparative vertebrate genomics field (Meadows & Lindblad-Toh, 2017), (Clark, Balciunas, et al., 2011).

The initial pGBT-RP 2.1 (RP2.1) vector has several features that efficiency cooperate to report gene sequence, expression and function (Clark, Balciunas, et al., 2011). Two main reporter components include a 5' protein trap and a 3' exon trap, with the entire cassette flanked by inverted terminal repeats (ITR) of the mini*Tol2* transposon to effectively deliver the transgene as single copy integrations into the zebrafish genome. In cases where RP2 integrates in the sense orientation of a transcription unit, the protein trap's splice acceptor overrides normal splicing of the transcription unit, creating a fusion between endogenous upstream exons and the monomeric RFP (mRFP) reporter sequences. The protein-trap domain in RP2.1 generates the expression profile, including subsequent protein localization and

accumulation when a functional in-frame fusion between the start codon-deficient mRFP reporter and the tagged protein. Mutagenesis is accomplished by the strong internal polyadenylation and putative border element, effectively truncating the endogenously tagged protein. The GBT mutagenesis system represented the first step toward a 'codex' of protein expression and functional annotation of the vertebrate genome (Clark, Balciunas, et al., 2011).

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We report here the development of a series of GBT protein trap vectors including versions to trap expression in each of the three potential reading frames. In addition, we modified the 3' exon trap to use a localized BFP rather than the more commonly used GFP to more effectively use these lines in conjunction with other transgenic fish. We deployed these vectors at scale, generating over 1000 protein trap lines with visible mRFP expression at either 2dpf (end of embryogenesis) or 4dpf (larval stage), with 36% and 87% of the cloned lines showcasing to our knowledge the first described expression of these genes at these stages, respectively. We used forward and reverse genetic tests to assess the mutagenicity of these vectors, noting similar rates of visible phenotypes at 5dpf as ENU and retroviral screening tools. We re-isolated five previously described loci, and embryos homozygous for each of the corresponding GBT alleles displayed strong loss of function phenotypes comparable to these previously published mutants generated using other mutagenesis strategies (ryr1b, fras1, tnnt2a, edar and hmcn1). Molecular assessment after positional cloning shows that nearly all alleles cause at least a 99+% knockdown of the tagged gene. Interestingly, over 35% of the cloned loci represent 68 mutants in zebrafish orthologs of human disease loci, including nervous, cardiovascular, endocrine, digestive, musculoskeletal, immune and integument systems. The GBT protein trapping system enabled the construction of a comprehensive protein codex including novel expression annotation, identifying new functional roles of the vertebrate genome and generating a diverse collection of potential models of human disease.

Materials and Methods

2 Zebrafish husbandry

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- 3 All zebrafish (*Danio rerio*) was maintained according to the guidelines and the standard procedures
- 4 established by the Mayo Clinic Institutional Animal Care and Use Committee (Mayo IACUC). The
- 5 Mayo IACUC approved all protocols involving live vertebrate animals (A23107, A21710 and A34513).
- 6 Generating GBT constructs, RP2 and RP8 series
- 7 pGBT-RP8.2 and -RP8.3 were made by combining three restriction endonuclease fragments of pGBT-
- 8 RP8.1, a 2.2 kb AfIII to AgeI, a 0.7 kb EcoRI to SpeI, and a 3.0kb SpeI to AfIII, with a short adapter to
- 9 close the space between AgeI and EcoRI that effectively removed one or two thymine nucleotides just
- 10 following the splice acceptor prior to the AUG-less mRFP cassette. For pGBT-RP8.2, Adapter-
- 11 GBT(+2) was made by annealing oligos adapter-GBT(+2)-a
- 12 [CCGGTTTTCTCATTCATTTACAGTCAGCCGG] and adapter-GBT (+2)-b
- 13 [AATTCCGGCTGACTGTAAATGAATGAGAAAA]. For pGBT-RP8.3, Adapter-GBT(+3) was made
- by annealing oligos adapter-GBT (+3)-a [CCGGTTTTCTCATTCATTTACAGCAGCCGG] and
- adapter-GBT(+3)-b [AATTCCGGCTGTAAATGAATGAGAAAA].
- pGBT-RP2.2 and -RP2.3 were made by combining three restriction endonuclease fragments of pGBT-
- 18 RP2.1 (Clark, Balciunas, et al., 2011), a 3.6kb BlpI to AgeI, a 1.9kb EcoRI to AvrII, and a 3.55kb AvrII
- to BlpI, with a short adapter to close the space between AgeI and EcoRI that effectively removed one or
- two thymine nucleotides just following the splice acceptor prior to the AUG-less mRFP cassette. For
- 21 pGBT-RP2.2, Adapter-GBT(+2) was made by annealing oligos adapter-GBT(+2)-a and adapter-GBT
- 22 (+2)-b. For pGBT-RP2.3, Adapter-GBT(+3) was made by annealing oligos adapter-GBT (+3)-a and
- adapter-GBT(+3)-b.

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pGBT-RP8.1 was made by cloning a mini-intron derived from carp beta actin intron 1 into pGBT-RP7.1. The 234bp SalI to XhoI mini-intron fragment was isolated from pCR4-bactmIntron following digestion. The pGBT-RP7.1 plasmid was digested with XhoI so that the SalI to XhoI fragment was cloned between the gamma-crystallin promoter and nls tagBFP. pCR4-bactmIntron was made by removing a 1.1kb internal portion of the carp beta actin intron 1 by digestion of pCR4-bact I1 with BstBI and BssHII, followed by filling in 5' overhangs and ligating remaining vector fragment. pCR4-bact_I1 was cloning a PCR product containing the carp beta-actin intron into pCR4-TOPO (Invitrogen). The intron was amplified from pGBT-RP2.1 (Clark, Balciunas, et al., 2011) using MISCbact_exon-F1 [CAGCTAGTGCGGAATATCATCTGCC] and MISC-bact_intron-R1 [CTTCTCGAGGTGAATTCCGGCTGAACTGTA] primers. pGBT-RP7.1 was made by replacing a 501bp PstI to PstI fragment of pGBT-RP6.1 with a 480bp PstI to PstI fragment of pRP2.1. This changed the nucleotide sequence between the carp beta-actin splice acceptor to replicate the sequences in pGBT-RP2.1. pGBT-RP7.1 was never directly tested in zebrafish. pGBT-RP6.1 was made by flipping the internal trap cassette relative the Tol2 inverted terminal repeats in pGBT-RP5.1. To do this, pGBT-RP5.1 was cut with EcoRV and SmaI. The 2.27kb EcoRV to SmaI vector backbone fragment, which included the ITRs, was ligated to the 3.51kb EcoRV to SmaI trap

1 fragment. pGBT-RP6.1 was then selected based on the right ITR of Tol2 being in front of the RFP trap, 2 which is the same orientation of pGBT-RP2.1. 3 pGBT-RP5.1 was made by cloning a PCR product with the AUG-less mRFP into pre(-1)GBT-RP5.1. 4 5 The 698bp mRFP* PCR product was obtained by amplification of pGBT-R15 (Clark, Balciunas, et al., 2011) with CDS-mRFP*-F1 [AAGAATTCGAAGGTGCCTCCTCCGAGGATGTCATCAAGG] and 6 7 CDS-mRFP-R1 [AAACTAGTCTTAGGCTCCGGTGGAGTGGCGG]. Prior to cloning the PCR 8 mRFP* product was digested with EcoRI and SpeI to prepare the ends for subcloning into pre(-1)GBT-9 RP5.1 that was opened between the carp beta actin splice acceptor and the ocean pout terminator. 10 pre(-1)GBT-RP5.1 was made by cloning 1.2kb SpeI to AvrII fragment from pGBT-PX (Sivasubbu et al., 11 2006) that contained the ocean pout terminator into the SpeI site of pre(-2)GBT-RP5.1. The resulting 12 products were screened for the proper orientation of the ocean pout terminator relative to the carp beta 13 actin splice acceptor. 14 15 pre(-2)GBT-RP5.1 was made by inserting an expression cassette to make a 3' poly(A) trap that makes 16 17 blue lenses. A 1.15kb SpeI to BgIII fragment from pKTol2gC-nlsTagBFP was cloned into pre(-3)GBT-RP5.1 that had been cut with AvrII and BgIII. This moved the *Xenopus* gamma crystallin promoter 18 driving a nuclear-localized TagBFP in front of the carp beta actin splice donor within pre(-3)GBT-RP5.1 19

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to create a localized BFP poly(A) trap signal replacing the ubiquitous GFP signal that was in pGBT-

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RP2.1.

- 1 pre(-3)GBT-RP5.1 was made by cloning a 492bp XmaI to NheI scaffold fragment from pUC57-I-
- 2 SceI_loxP_splice into pKTol2-SE (Clark, Balciunas, et al., 2011) opened with XmaI and NheI.
- 3 pUC57-I-SceI_LoxP_Splice contains a synthetic sequence (see below) cloned into pUC57 (Genscript).
- 4 The scaffold contains an I-SceI site; loxP site; carp beta actin splice acceptor; cloning sites for mRFP,
- 5 ocean pout terminator, and BFP lens cassettes; carp beta actin splice donor; loxP site; and an I-SceI site.
- 6 [cccgggatagggataacagggtaatataacttcgtatagcatacattatacgaagttatcgttaccaccactagcggtcagactgcagattgcagcac
- 8 gcctgttacctgcactcaccgacaagctgttaccctggaattcgtttaaacactagtcaccggcgttcctaggttataagatctacctaaggtgagttgatct
- 10 catttacactgagctcaagacgtctgataacttcgtatagcatacattatacgaagttattaccctgttatccctatggctagc]

Generating GBT collection

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- Generation of the GBT collection was based on the prior described protocols (Clark, Balciunas, et al.,
- 14 2011; Clark, Urban, Skuster, & Ekker, 2011; J. Ni et al., 2016).

Fluorescent microscopy of mRFP reporter protein expression

- Larvae were treated with 0.2 mM phenylthiocarbamide at 1 dpf to inhibit pigment formation. The
- anesthetized fish were mounted in 1.5% agarose (Fisher Scientific BP1360) prepared with 0.017mg/ml
- 19 tricaine solution in an agarose column in the imaging chamber. The protocol of ApoTome microscopy
- was described in previous publication. (Clark, Balciunas, et al., 2011) For Lightsheet microscopy, larval
- 21 zebrafish were anesthetized with 0.017g/ml tricaine (Ethyl 3-aminobenzoate methanesulfonate salt) in
- 22 embryo water during imaging procedure. To capture RFP expression patterns of 2 dpf and 4 dpf larval
- 23 zebrafish, LP 560 nm filter as excitation and LP 585nm as emission was used for Lightsheet microscopy.

- 1 The sagittal-, dorsal-, and ventral- oriented z-stacks of the mRFP expression were captured at either 50x
- 2 magnification using an ApoTome microscope (Zeiss) with a 5x/0.25 NA dry objective (Zeiss) or 50x
- 3 magnification using a Lightsheet Z.1 microscope (Zeiss) 5x/0.16 NA dry objective. Each set of images
- 4 were obtained from the same larva and the images shown are composites of the maximum image
- 5 projections of the z-stacks obtained from each direction.

Sperm Cryopreservation

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- 8 Sperm collection and cryopreservation was initially based on the protocol described in (Draper & Moens,
- 9 2009) and moved to the ZIRC protocol described in (Matthews et al., 2018).

Genomic DNA isolation

- Genomic DNA was isolated from F1 fish tail biopsies to conduct next generation sequencing and from
- both WT and heterozygous larva to manually perform the PCR-based mRFP linkage analysis. Zebrafish
- larva were individually placed to 0.2 ml PCR tubes and sacrificed to extract genomic DNA in 50 mM
- 15 NaOH for 20 min at 95 C°.

PCR-based linkage analysis of GBT insertions loci

- 18 TALE-PCR: The protocol used was designed to amplify and clone junction fragments from Tol2-based
- 19 gene-break transposons (GBT) in the zebrafish genome. Although modified, it is based on a protocols
- 20 received from Alexi Parnov, Vladimir Korsch, and Karuna Sampath. The following primer mixtures
- 21 (containing 0.4 μM GBT specific primer and 2 μM DP primer) were prepared: for primary PCR: 5R-
- 22 mRFP-P1/DP1, 5R-mRFP-P1/DP2, 5R-mRFP-P1/DP3, 5R-mRFP-P1/DP4, 3R-GM2-P1/DP1, 3R-
- 23 GM2-P1/DP2, 3R-GM2-P1/DP3, 3R-GM2-P1/DP4, 3R-tagBFP-P1/DP1, 3R-tagBFP-P1/DP2, 3R-

- tagBFP-P1/DP3, 3R-tagBFP-P1/DP4; for secondary PCR: 5R-mRFP-P2/DP1, 5R-mRFP-P2/DP2, 5R-
- 2 mRFP-P2/DP3, 5R-mRFP-P2/DP4, 3R-GM2-P2/DP1, 3R-GM2-P2/DP2, 3R-GM2-P2/DP3, 3R-GM2-
- 3 P2/DP4, 3R-tagBFP-P2/DP1, 3R-tagBFP-P2/DP2, 3R-tagBFP-P2/DP3, 3R-tagBFP-P2/DP4; for tertiary
- 4 PCR: TAIL-bA-SA/DP1, TAIL-bA-SA/DP2, TAIL-bA-SA/DP3, TAIL-bA-SA/DP4, Tol2-ITR(L)-
- 5 O1/DP1, Tol2-ITR(L)-O1/DP2, Tol2-ITR(L)-O1/DP3, Tol2-ITR(L)-O1/DP4, Tol2-ITR(L)-O3/DP1,
- 6 Tol2-ITR(L)-O3/DP2, Tol2-ITR(L)-O3/DP3, Tol2-ITR(L)-O3/DP4. A total of 1 μl of primer mixtures
- 7 were added to PCR reaction (total volume 25 μl). Cycle settings were as follows. Primary: (1) 95°C, 3
- 8 min; (2) 95°C, 20 sec; (3) 61°C, 30 sec; (4) 70°C, 3 min; (5) go to "cycle 2" 5 times; (6) 95°C, 20 sec;
- 9 (7) 25°C, 3 min; (8) ramping 0.3°/sec to 70°C; (9) 70°C, 3 min; (10) 95°C, 20 sec; (11) 61°C, 30 sec;
- 10 (12) 70°C, 3 min; (13) 95°C, 20 sec; (14) 61°C, 30 sec; (15) 70°C, 3 min; (16) 95°C, 20 sec; (17) 44°C,
- 11 1 min; (18) 70°C, 3 min; (19) go to "cycle 10" 15 times; (20) 70°C, 5 min; Soak at 12 °C. A total of 5
- 12 μl of the primary reaction was diluted with 95 μl of 10mM Tris-Cl or TE buffers and 1μl of the mixture
- was added to the secondary reaction. Secondary: (1) 95°C, 2 min (2) 95°C, 20 sec; (3)61°C, 30 sec; (4)
- 70°C, 3 min; (5) 95°C, 20 sec; (6) 61°C, 30 sec; (7) 70°C, 3 min; (8) 95°C, 20 sec; (9) 44°C, 1 min;
- 15 (10) ramping 1.5°/sec to 70°C; (11) 70°C, 3 min; (12) go to "cycle 2" 15times; (13) 70°C, 5 min; Soak
- at 12°C. A total of 5 µl of the primary reaction was diluted with 95 µl of 10mM Tris-Cl or TE buffers
- and 1µl of the mixture was added to the tertiary reaction. Tertiary: (1) 95°C, 2 min; (2) 95°C, 20 sec; (3)
- 18 44°C, 1 min; (3) ramping 1.5°/sec to 70°C; (4) 70°C, 3 min; (5) go to "cycle 2" 32 times; (6) 70°C, 5
- min; Soak at 12°C. Products of the secondary and tertiary reactions were separated by using 1-1.5%
- agarose gel. The individual bands from the "band shift" pairs were cut from the gel and purified by
- 21 using QIAquick Gel Extraction Kit (QIAGEN, Germany), and sequenced by using ABI Cycle
- 22 Sequencing chemistry (PE Applied Biosystems, CA) and an ABI Prism 310 Genetic Analyzer with Data
- 23 Collection Software (PE Applied Biosystems, Foster City, CA) supplied by the producer.

1 5' and 3' RACE PCRs: The protocol used was designed to use cDNA to amplify and clone junction 2 fragments of Tol2-based gene-break transposons (GBT) in the zebrafish genome. 5' RACE allows PCR 3 amplification of unknown sequence at the 5' end of a cDNA as long as there is enough known sequence 4 within the cDNA to design two antisense primers. Although modified, it is based on the protocol in described in (Clark, Balciunas, et al., 2011). The following primer mixtures were prepared: for primary 5 PCR: 0.20 µM GBT specific primer (5R-mRFP-P1), and a mix of universal 5' RACE primers 2.5 µM 6 7 5R-UP-S and 0.5 μM 5R-UP-L. Secondary reaction: 25 μM GSP (5R-mRFP-P2), and 25 μM universal 8 primer 5R-N1. The reaction mix used was as follows: Primary: (25 µl reaction) Template (RR-cDNA) 2 μl, Bioline 5X myTag buffer 5 μl, myTag 0.25 μl, GSP 0.5 μl, Universal 5' RACE primer mix (URS 9 mix) 2.0 μl, Water 15.25 μl. Secondary: (50 μl reaction) Template (2:100 dilution 1° PCR Reaction), 10 Bioline 5X myTaq buffer 10 μl, myTaq 0.3 μl, GSP-P2 0.9 μl, URS-P2 0.9 μl, Water 35.9 μl. Cycle 11 12 Settings are as follows. Primary: (1) 95°, 3'; (2) 95°, 30"; (3) 65°, 30" -0.5°/cycle; (4) 70°, 2'; Go To (2) x 15 cycles; (5) 95°, 30"; (6) 57°, 30"; (7) 70°, 2'; Go To (5) x 20 cycles, (8) 70°, 10'; (9) Soak at 12°. 13 Dilute 2µL of the primary PCR reaction with 198µL of 10mM Tris-Cl; 1mM EDTA pH8.0. Secondary: 14 15 (1) 95° for 3' (2) 95°, 30"; (3) 63°, 30" -0.5°/cycle; (4) 70°, 2'; Go To (2) x 10 cycles; (5) 95°, 30"; (6) 58°, 30"; (7) 70°, 2'; Go To (5) x 25 cycles; (8) 70°, 10'; Soak at 12°. After the PCR reactions finish 16 17 20µL of each sample were run on a 1.2% agarose gel. You may also run 10µL of undiluted primary PCR reactions on the same gel; however, most of the time the bands of interest do not all appear until 18 after the nested PCR reactions have been run. The individual bands from the gel were excised and 19 20 purified by using QIAquick Gel Extraction Kit (QIAGEN, Germany), and sequenced by using ABI Cycle Sequencing chemistry (PE Applied Biosystems, CA) and an ABI Prism 310 Genetic Analyzer 21 22 with Data Collection Software (PE Applied Biosystems, Foster City, CA) supplied by the producer.

Forward genetic screening with next-generation sequencing

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- 2 Isolated genomic DNA (300-500ng) was digested with MseI, and BfaI in parallel for 3h at 37°C and
- 3 heat inactivated for 10 min at 80°C. The digested samples from each enzyme were pooled with
- 4 prealiquoted barcoded linker in individual wells. The T4 DNA ligase (New England Biolabs, Inc.) was
- 5 added, and the reaction mix was incubated for 2 h at 16°C. The linker-mediated PCR was performed in
- 6 two steps. In the first step, PCR was done with one primer specific to the 3'- ITR (5'-
- 7 GACTTGTGGTCTCGCTGTTCCTTGG-3') and the other primer specific to linker sequences (5'-
- 8 GTAATACGACTCACTATAGGGC- 3') using the following conditions: 2 min at 95°C, 25 cycles of 15
- 9 sec at 95°C, 30 seconds at 55°C and 30 seconds at 72°C. The PCR products were diluted to 1:50 in
- 10 dH2O, and a second round of PCR was performed using ITR (5'-
- 11 TCACTTGAGTAAAATTTTTGAGTACTTTTTACACCTC-3') and linker specific (5' -
- 12 GCGTGGTCGACTGCGCAT-3') nested primers to increase sensitivity and avoid non-specific
- amplification using the following conditions: 2 min at 95°C, 20 cycles of 15 sec at 95°C, 30 seconds at
- 14 58°C and 30 seconds at 72°C. The nested PCR products from each 96-well plate are pooled and
- processed for Illumina library preparation as per manufacturer's instructions.

Protein classification of the cloned zebrafish genes

- The 183 cloned zebrafish genes are classified by using PANTHER (Mi, Muruganujan, & Thomas, 2013).
- 19 PANTHER provided protein classes of the molecule coded by the cloned zebrafish genes.

Annotating human orthologues of GBT-tagged genes and disease-causing genes

- 22 The human orthologues of 192 cloned zebrafish genes were mainly collected by using a data mining tool,
- 23 ZebrafishMine (Van Slyke et al., 2018) supported by the ZFIN database. In some cases, the candidates

of human orthologues unlisted in ZFIN database were manually searched by using both Ensembl

2 (https://useast.ensembl.org/index.html) and InParanoid8 (https://inparanoid.sbc.su.se/cgi-bin/index.cgi)

databases. In parallel, the candidates were manually identified by the result of BLASTP assembled with

4 human proteins and by the result of an online synteny analysis tool, SynFind

5 (https://genomevolution.org/CoGe/SynFind.pl). If the candidate multiply hit in those manual

assessments, it was annotated as a human orthologue. The human phenotype data caused by mutations

of 68 human orthologues were collected by using another data mining tool, BioMart

(http://useast.ensembl.org/biomart/martview/cfe15ead83199a0b7c7997f5a4ce9e6b) supported by

9 Ensembl database.

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Finding Disease Models in Vertebrates

- Mouse models were found by using both descriptions of animal models in Online Mendelian Inheritance
- in Man (OMIM; https://www.omim.org/) and in Mouse Genome Informatics (MGI;
- 14 http://www.informatics.jax.org/humanDisease.shtml). MGI provided the details of mouse models of
- 15 human disease, such as the number of models have been established. Zebrafish model were also found
- by using both OMIM and the Zebrafish Information Network (ZFIN; https://zfin.org/). ZFIN provided
- all data of fish strains listed in this database.

Gene expression profiling of the cloned zebrafish genes

20 The cloned genes with unpublished expression data were isolated by using "Gene Expression" tool of

ZebrafishMine (Van Slyke et al., 2018). In parallel, some published expression data were also manually

searched from ZFIN database or in some references. To isolate the gene with the expression localized in

the tissues or the organs shows abnormalities in the causing diseases of the human orthologues, the

- 1 mRFP reporter expression patterns of the cloned genes 2 and 4 dpf are manually analyzed using
- 2 zfishbook database (Clark, Argue, Petzold, & Ekker, 2012).

Results

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The features of GBT constructs RP2 and RP8 – capturing all three proteomic reading frames 2 3 In our previous study, we reported the intronic-based gene-breaking transposons (GBTs) as effective and revertible loss-of function tools for zebrafish (Clark, Balciunas, et al., 2011). The main features of the 4 RP2.1 vector system are as follows (Fig1A): 1) Genetically engineered cargo is flanked by miniTol2 6 sequences necessary and sufficient for Tol2 transposase-mediated transposition, an efficient transgenesis 7 vector in zebrafish (Kawakami et al., 2004); (Balciunas et al., 2006); (Urasaki, Morvan, & Kawakami, 2006); 2) protein trap that enables in vivo expression selection of the vertebrate proteome. The addition 8 9 of the AUG-free mRFP reporter has yielded an effective protein trap for both organ-specific and 10 subcellular localization of the tagged locus (Petzold et al., 2009); (Clark, Balciunas, et al., 2011); (Liao 11 et al., 2012);(Xu et al., 2012);(Ding et al., 2013); (Westcot et al., 2015); 3) Mutagenic transcriptional terminator. The 5' cassette is a combination of a strong splice-acceptor (SA), poly adenylation signal 12 (pA), and putative border element (red octagon) in conjunction with a start codon (AUG)-free monomeric red fluorescent protein (mRFP) reporter. These elements have been shown to be effective as a transcriptional stop in zebrafish by hijacking endogenous splicing (Sivasubbu et al., 2006). These elements are very effective at inducing a quantitative knockdown in all 26 lines assessed to date using qRT-PCR; 97% or higher knockdown in all lines (Clark, Balciunas, et al., 2011); (Ding et al., 2013); (Ding et al., 2016), this manuscript). The GBT mutagenesis system is thus an effective first step to 18 creating a gene codex simultaneously combining expression and loss of function genetics. 20 However, some limitations were noted with the initial RP2.1 vector – notably the effective trapping of transcripts without detectable expression of the mRFP reporter. Molecular cloning of these GFP+/RFP-22 lines demonstrated the fidelity of expression requiring the capture of an appropriate reading frame.

RP2.1 was designed to use one main reading frame, and some lines with expression were noted to

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include the use of a secondary splice acceptor (data not shown). To maximize genome coverage of this insertional mutagen, we created all three reading frames of the RP2 and RP8 vector series (Fig. 1). 4) 3' exon trap. These vectors also encodes a 3' exon trap with preferential expression following intragenic insertions in zebrafish (Sivasubbu et al., 2006); (Petzold et al., 2009); (Clark, Balciunas, et al., 2011). The function of this cassette complements the obligate, in-frame protein trapping effect and is used for both quality control during mutagenesis and for genotyping of more weakly expressing protein trap alleles (Fig.1A). In the RP2 vector series, the nearly ubiquitous b-actin promoter drives expression of GFP. Expression of integrated GFP becomes detectable between early developmental stages such as seven- to eight-somite-stage and provides bright expression with a good signal-to-noise ratio at 25 hpf (Davidson AE et al., 2003). However, the ubiquitous GFP expression from the 3' exon trap cassettes could interfere with another fluorescent marker system based on GFP labeling in further studies. The RP8 vector series includes all reading frames for the AUG-free mRFP reporter and a new 3' exon trap cassette with expression of tagBFP driven by the lens-specific gamma-crystalline promoter (Fig. 1B). Using the tissue-specific reporter system with BFP is helpful to easily detect F1 founder with weak mRFP expression and to avoid interference with GFP-based multi-labeling purposes when crossed with other GFP-labeled transgenic fish lines. 5) Revertible mutagenic cassette. The flanking loxP sites enable reversion of the tagged locus by Cre – mediated recombination. This facilitates both somatic (Clark, Balciunas, et al., 2011); (Ding et al., 2013) and germline approaches (Petzold et al., 2009). We generated more than eleven hundred independent lines by using all six constructs of the GBT system (Supplemental Table 1). We conducted an initial screening expression of the mRFP fusion protein and showed that RP2 and RP8 vector series with all reading frames of mRFP reporter protein readily detects the distribution of the fusion proteins expressed from their own promoter in zebrafish (Supplemental Figure 1).

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Annotation of protein localization and trafficking of the GBT strain collection The ability to non-invasively obtain temporal and spatial expression pattern information is a key feature of these protein trap strains. Pilot data from our first RFP lines rapidly demonstrated this step was going to be a major bottleneck for our pipeline if we used standard documentation methods. Consequently, we established a capillary-based confinement and imaging protocol (SCORE imaging; (Petzold et al., 2010) for quickly capturing and holding living zebrafish for rapid, high quality fluorescent expression in precise longitudinal imaging angles. The animals are easily inserted into a capillary of the correct diameter for the particular fish stage, then placed on the microscope and covered with a matched solution to remove the optical distortion from the capillary housing. The capillary is simply rotated for precise 0 degree (dorsal), 90 degree (lateral), and 180 degree (ventral) images on standard fluorescent microscopes, such as the Apotome. The technical bottleneck of standard microscopy to scan a whole embryo and larva is also labor-intensive and time-consuming. For instance, fluorescent scanning of a quarter of a 2dpf or 4dpf larvae required about 15 min exposure using a Zeiss Apotome microscope, one imaging modality deployed for this collection. To accelerate the expression profiling of these GBT lines, we subsequently utilized a Zeiss Lightsheet Z.1 SPIM microscope. The Lightsheet enabled high speed scanning of a whole embryo or larva, resulting in a nearly 20x faster image acquisition rate than the primary imaging process utilizing the Apotome. We prioritized and cataloged lines with robust expression at 2 and 4 days post fertilization (dpf). All zebrafish lines are freely available now through zfishbook (Clark et al., 2012) and are partially accessible from the Zebrafish International Resource Center. Consequently, we have generated 1138 lines by using each vector system showing in Supplemental Table 1), and updated results are posted at zfishbook.

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Our throughput for cloning the GBT lines using traditional molecular methods was clearly an initial bottleneck. To help address this, we deployed a rapid cloning process based on methods used to isolate retroviral integrations (Varshney, Huang, et al., 2013; Varshney, Lu, et al., 2013). This method leverages the massive parallel sequencing technology of the Illumina MiSeq, yielding 101 bp sequencing reads, followed by a custom bioinformatics pipeline that involves both mapping and annotation. Fin-clips from four male animals per GBT locus are obtained during sperm cryopreservation and are used as a source of DNA. Shared inserts in multiple individuals from a single GBT line are considered candidate loci. This information is subsequently used to generate gene-specific primers to manually confirm linkage using 5° or 3'RACE, inverse PCR, or TAIL PCR to generate locus-specific primers for downstream molecular genotyping applications. 212 of the lines met the highest stringency of confirmed expression linkage and are classified as confirmed GBT integrations. An additional 143 lines have been initially tagged using this high throughput method, yielding candidate integration annotation as listed in zfishbook. The ability to complete the annotation status from candidate to confirmed for any given GBT locus with a desired expression profile is enhanced by the continued refinement of the zebrafish genome. mRFP Expression profiling reveals overlap with known annotation at 2dpf and substantive new expression data at both 2dpf and 4dpf GBT lines are currently imaged to capture mRFP expression at both 2 and 4 dpf, including dorsal, sagittal and ventral views using the SCORE imaging method. (Petzold et al., 2010) In an openly accessible database of GBT lines, zfishbook, the images of mRFP expression pattern were stored within the media gallery associated with each line (Clark et al., 2012). We summarized published expression data of the tagged genes in both zfishbook and ZFIN in Table 1. Imaging the localization of transcripts

and proteins at 4dpf is more difficult than those at 2 dpf, because accessibility of antisense RNA probes

and antibodies into the larva's body is technically limited for in the methods of both in situ hybridization

and immunohistochemistry. Compared with the published data of gene expression in ZFIN, zfishbook

currently provides almost the double number of genes with expression data at 2 dpf and 14 times the

number of genes with expression data at 4 dpf (Table 1). In addition, zfishbook also provides novel

expression data for 61 genes at any developmental stage (Fig.2).

High knockdown efficiency of endogenous transcripts induced by RP2

9 We directly compared published transposon insertional mutant vector systems (Fig. 3). The range and

average knockdown levels in the FlipTrap system (Trinh le et al., 2011) produced a range of 4-30% (70-

96% knockdown capacity) in six tested fish alleles, a similar range to our initial R-series protein trap

vectors (R14-R15) that used a simple transcriptional terminator (Liao et al., 2012; Petzold et al., 2009)

The pFT1 appears to be an improvement over these systems, in which the overall range and average

read-through is reduced to 6-11% (89-94% knockdown) from four tested fish alleles (T. T. Ni et al.,

2012). In contrast, the RP2.1 vector (Clark, Balciunas, et al., 2011), (Ding et al., 2013), (Ding et al.,

2016) and this manuscript) maintains a strong knockdown (1% or less read-through) in 26 lines tested.

Though deployed here using a nearly random, transposon-based delivery platform, the GBT vector

system is an effective insertional mutagen suitable for an array of other – including targeted integration -

genome-wide applications.

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Phenotypic Appearance Rate is Similar to Other Mutagenic Technologies for Forward Genetics

Screening through 5 dpf

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We conducted an initial forward genetic screen on embryos and early larvae of 179 RFP-positive GBT lines, identifying 12 recessive phenotypes, such as ryr1b, fras1, tnnt2a, edar and hmcn1, (Clark, Balciunas, et al., 2011; Westcot et al., 2015) visible during the first five days of development including lethality, heart, muscle, skin and other phenotypes. This 7% recovery of visible early developmental mutants is very similar to the 5% recovered visible mutants from the Sanger TILLING consortium analysis of truncated zebrafish genes (Kettleborough et al., 2013). This 7% is also comparable to prior retroviral (Amsterdam & Hopkins, 2004) and ENU(Haffter et al., 1996) zebrafish mutagenesis work that estimated between 1400 and 2400 genes (~5-9% of the genome) would result in a visible embryonic phenotype when mutated. **GBT** alleles phenocopy known embryonic mutations We tested the first five GBT lines in genes with known loss of function mutant phenotypes (ryr1b; fras1; tnnt2a; edar; hmcn1). All five of these alleles in genes with described loss of function defects are phenocopied by these GBT insertional alleles (Clark, Balciunas, et al., 2011); (Westcot et al., 2015); this manuscript). These loci represent a critical internal methods reference further validating the mutagenicity of these novel insertional vectors. Gene ontology analysis of GBT-tagged loci To assess the diversity of GBT loci molecularly characterized to date, we utilized the PANTHER classification system (v.14.0, pantherdb.org, (Mi et al., 2013) and generated a table of protein class ontology tags in the molecularly isolated GBT lines. The PANTHER Protein Class ontology was adapted from the PANTHER INDEX (PANTHER/X) ontology that comprises two types of classifications: molecular function and biological process and includes commonly used classes of

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protein families. The molecular function schema classifies a protein based on its biochemical properties, such as receptor, cell adhesion molecule, or kinase. The biological process schema classifies a protein based on the cellular role or process in which it is involved, for example, carbohydrate metabolism (cellular role), TCA cycle (pathway), neuronal activities (process), or developmental processes (process) (Thomas PD et al, 2013). As of April 2018, almost half of known zebrafish genes (10626/25289 genes) were tagged in the PANTHER Protein Class ontology. 168 of our cloned GBT alleles mapped in the PANTHER system with 21 types of Protein Classes (Table 2). 18% and 16 % of the mapped GBT alleles are classified to nucleic acid binding (PC00171) and transcription factor (PC00218), respectively (Fig. 4). This result reveals that a quarter of the mapped genes possibly play a role in regulatory processes. Overall, however, the rich diversity of protein classes observed in our cloned traps suggests a large diversity will be represented by the overall collection and consistent with the random nature of genome integration events by the Tol2 transposon (Clark, Balciunas, et al., 2011). Disease-causing human orthologues of GBT-tagged genes Of 183 GBT-tagged genes, 171 human orthologues were annotated in at least one public database such as ZFIN, Ensembl, Homologene, and InParanoid (Table 3). Several human orthologues were provisionally annotated using BLASTP and a synteny analysis tool, SynFind. In a previous study comparing the list of human genes possessing at least one zebrafish orthologue with the 3,176 genes bearing morbidity descriptions that are listed in the OMIM database, 82 % morbid genes 2,601 genes) can be related to at least one zebrafish orthologue (Howe et al., 2013). Surprisingly, 67 genes (about 37%) of 183 annotated human orthologues are associated with human disease involved in multi-organ system including nervous, circulatory, endocrine, metabolic, digestive, musculoskeletal, immune, and integument systems (Fig. 5 and Table 3) and many are not established in rodents and zebrafish (Table

- 5). The GBT protein-trap system provides a variety of potential human disease models which have a
- 2 revertible allele that can interchange between disease and healthy cellular, organ and physiological states.
- 4 The GBT Protein Trap Reveals Protein-Coding Regions Not Predicted by the Zebrafish Genome
- 5 **Project**

- 6 16 of 211 molecularly confirmed lines by manual, PCR-based mRFP linkage analysis using TALE,
- 7 inverse, and RACE PCRs (Table 4) do not match any expressed sequence tag (EST) or predicted genes.
- 8 However, in each case we were able to confirm transcription at the locus in wild-type animals, yielding
- 9 new annotation for these loci in the zebrafish genome.

The RP2 and RP8 vectors of GBT system were assembled to capture all three reading frames of fusion

Discussions

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Novel Expression Annotation Revealed By Protein Trapping of Endogenous Genes

protein of the trapped gene with the mRFP reporter. This GBT system reveals that mRFP-truncated 4 5 fusion proteins exhibit distinct subcellular localization. This is particularly noteworthy for the 4dpf stages because published gene expression data at late developmental stages have been limited by the 6 7 technical difficulties in conducting such analyses using traditional techniques such as whole mount in situ hybridization (WISH) at these larval and later time points. We note that the mRFP-truncated fusion 8 9 protein may localize ectopically in cases where the protein localization signal is contained in the Cterminal domain (Clark, Balciunas, et al., 2011; Trinh le & Fraser, 2013). Although the extent that 10 11 subcellular localization recapitulates the endogenous protein is dependent on each insertion locus specifics, the visualizing and illuminating spatiotemporal expression patterns of trapped protein may 12 13 facilitate dynamic studies of specific cell types and molecular functions in a living vertebrate. The novel

expression description at 4dpf in nearly five of six cloned loci demonstrates the dearth of 3D expression

annotation for the overwhelming number of genes, even in one of the most studied model system such as

the zebrafish. With ever-improving microscope-based imaging tools (Liu et al., 2018), these lines have

the potential to help annotate at diverse developmental and adult stages, while also potentially imaging

Gene-Break Protein Trap in an Effective Insertional Mutagen with High Knockdown Efficiency

and Cre-Reversion to WT allele

subcellular expression for a subset of protein trap fusions.

1 Transposons offer several unique features over and above traditional static mutational approaches, 2 including high quality expression tools and new regulated mutagenesis methodologies. From the 3 perspective of genome engineering development, GBT technology was the first method for revertible 4 allele generation of vertebrates outside of the mouse (Clark, Balciunas, et al., 2011); (Ding et al., 2013). 5 We know two major potential biases that may yield non-random trapping coverage of the genome. First, 6 the RP2.1 protein trap was initially designed around a single reading frame. Upon molecular analysis of 7 our first lines, however, we discovered that RP2.1 encodes a second, alternative splice acceptor yielding protein trap expression from a second reading frame due to this alternative splicing event in a significant 8 9 number of our lines. The deliberate development of RP2 and RP8 vectors for each reading frame 10 obviates this potential limitation when used in other delivery modalities besides the Tol2 transposon. For example, these vectors would be suitable for gene editing-based targeted knockin methods. 11 The GBT system deployed here has been joined by two new and complementary transposon 12 13 mutagenesis systems. The FlipTrap system by Dr. Fraser's group(Trinh le et al., 2011), which can be mutagenic when provided Cre recombinase, is primarily focused on imaging fusion proteins in vivo and 14 addressing cellular dynamics and related questions. The FT1 system by the Chen lab is a complementary flipping trap that can use either Cre or Flp recombinases to regulate alleles depend on the original orientation of the insertion (T. T. Ni et al., 2012). GBT-based zebrafish alleles are highly 18 complementary and non-redundant to those generated by other mutagenesis methods, including these 19 other transposons and demonstrated higher knockdown efficiency of the WT transcripts than those mutagens because of the use of an enhanced polyadenylation signal and a putative boundary element 20 21 between 5' protein trap and 3' exon trap cassettes. Importantly, since the initiation of this project to 22 generate a collection of Cre-revertible mutant alleles, several groups have now reported collections of tissue-specific Cre driver lines including the Brand lab (Jungke et al., 2013; http://crezoo.crt-

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dresden.de/crezoo/), the Zcre consortium (http://zcre.org.uk/) and the Wen lab in the PTC Consortium .

The GBT system described here is a two-component, molecularly regulatable mutagenesis approach that

offers the ability to test for the sufficiency of protein-encoding loci in regulated, tissue- and cell- specific

applications.

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Functional Diversity of the Trapped Proteins by the GBT system

To analyze distribution of protein functions of the trapped genes by GBT system, we performed GO

analysis using the PANTHER protein classification. Although the protein functions related in

transcriptional regulatory process, such as nucleic acid binding and transcription factors represented one

relatively common class of isolated genes, the protein GO analysis indicated that the GBT protein trap

was a useful tool for capturing a wide range of protein functions in addition to cell fate regulators and

related nuclear genes.

Cloned GBT Loci Represent a Rich Collection of Potential Human Disease Models

More than 7,000 human diseases have already been described, and 80% of those are thought to have a

genetic origin (Varga et al., 2018). Model organism studies can be a pivotal resource for understanding

gene function which possibly provides additional insight into the cause of particular disease, thereby

contributing to understanding of the pathogenic process and discovery of the therapeutic strategy

(Wangler et al., 2017). Annotation of human orthologues of 160 tagged genes revealed that GBT

technology yield a high frequency potential human disease models with loss of gene function, tracking

expression of the truncated protein and Cre-revertible mutated allele to rescue the phenotype.

Furthermore, the modeling human genetic diseases using the GBT system has advantages compared with reverse genetic approaches such as TALEN and CRISPR-Cas9 systems. In initial phenotype

with reverse genetic approaches such as TALEN and CRISPR-Cas9 systems. In initial phenotype

screening, PR2.1 mutagenesis demonstrated 7% phenotype appearance as much as the other ENU- and retroviral forward genetic screenings (Amsterdam & Hopkins, 2004; Haffter et al., 1996; Kettleborough

et al., 2013). For example, the annotation of disease-causing human orthologues of the tagged genes also

revealed that the GBT system comprehensively developed mutants in zebrafish orthologues of human

disease loci, including nervous, cardiovascular, endocrine, digestive, musculoskeletal, immune, and

integument systems. Surprisingly, this system generated 68 of pioneering mutants in orthologs of human

disease loci (Supplemental Table 2).

Discovery of Novel Transcripts by Trapping Unpredicted Genes

Since the completion of the zebrafish reference genome sequencing, it has enabled many new discoveries to be made, in particular the positional cloning of hundreds genes from mutation affecting embryogenesis behavior, physiology, and health and disease. However, a few poorly assembled regions remain (Howe et al., 2013). In molecular cloning of GBT lines generated, we found that a surprising proportion of the sequenced insertions does not correspond to any predicted genes. Although we have not formally excluded that mRFP expression might, in some case, be an artifact, the data of gene prediction provided in genome databases reveals some prediction errors. These results suggest that the algorithms used to predict genes from genome databases have missed a significant number of genes. The protein trapping by using GBT system may useful in identifying unsuspected novel genes, expressions and functions *in vivo* in real time.

New Genomic Insights Using the GBT Random Insertion Mutagen

to the use of the zebrafish to annotate the vertebrate genome.

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2 The ready ease of these mRFP-based protein traps for basic expression analyses demonstrates how much 3 we do not know about our overall proteome and the codex that is our genome. Nearly forty percent and 4 five of six of the cloned genes show no expression at 2dpf and 4dpf, respectively, in ZFIN. At the 5 subcellular level, such protein traps in conjunction with new microscopy techniques represent just 6 another method for cellular and mechanistic analyses in an in vivo cellular context. Although these were 7 made using random insertional approaches, new targeted integration tools using gene editing such as GeneWeld(Wierson et al., 2018) should readily empower labs to build their custom GBT lines in the 8 9 future for genes not in this collection. Together, this initial 1100+ GBT collection is a new contribution

1 Competing interests

2 The authors declare no competing interests.

- 1 Figure legends
- 2 Table 1 Comparison availability of expression data
- 3 Table 2 Protein classification
- 4 Table 3 Disease-causing human orthologues
- 5 Table 4 Potential novel human disease models
- **6 Table 5 Novel transcripts**
- 7 Figure 1 Schematic of the RP2 and RP8 gene-break transposon system with all three reading
- 8 frames of AUG-less mRFP reporter
- 9 A. Schematic of the RP2 system fused with 3 reading frames of AUG-less mRFP reporter (RP2.1, RP2.2
- and RP2.3). B. Schematic of the RP2 system fused with 3 reading frames of AUG-less mRFP reporter
- 11 (RP8.1, RP8.2 and RP8.3). ITR, inverted terminal repeat; SA, loxP; Cre recombinase recognition
- sequence, splice acceptor; *mRFP' AUG-less mRFP sequence; poly (A)+, polyadenylation signal; red
- octagon, extra transcriptional terminator and putative border element; β -act, carp beta-actin enhancer,
- SD, splice donor; E, enhancer; P, promoter; and WT, wild type.
- 15 Figure 2 Novel protein expression
- 16 Representative novel expression data of trapped proteins fused to the mRFP reporter in this GBT
- 17 collection. A. unkl localized in olfactory pit, cerebrum and spinal cord at 4dpf. B. nusap1 localized in
- retina and the top layer of both forebrain and midbrain at 4 dpf. C. zgc:194659 strongly expressed in the
- brain and spinal cord at 4 dpf. D. marcksl1a expressed in the lens, skin and notochord at 2 dpf. E. pipp2a
- specifically localized to otoliths at 2 dpf. F. ahnak specifically expressed in skin at 4 dpf. G. dph1
- 21 ubiquitously expressed showing granulized pattern in somites at 4 dpf. H. nfatc3a expressed in skeletal
- muscle and skin at 4 dpf. I. pard3bb localized to the pronephros and gut at 4 dpf. White arrowhead

- shows an artificial expression in lens driven by the promoter of 3' exon trap of RP8.1. J.
- 2 LOC100537272 expressed in circulatory cells in the blood stream at 4 dpf.
- 3 Figure 3 High knockdown efficiency of RP2.1 compared with other previous gene-trap systems
- 4 Black dots of bar graphs shows percentage of remaining endogenous transcripts in homozygous larvae
- 5 with mean and 95% confidence interval indicated by individual lines. The data of previous protein trap
- 6 systems were also converted from the data in the original articles, R14-R15, our initial R-series protein
- 7 trap vectors (n= 6),(Clark, Balciunas, et al., 2011); FlipTrap, FlipTrap vectors (n= 6), (Trinh le et al.,
- 8 2011); FT1, FT1 vector (n=4),(T. T. Ni et al., 2012); RP2.1 (n=26), (Clark, Balciunas, et al., 2011; Ding
- 9 et al., 2013; El-Rass et al., 2017; Westcot et al., 2015) and unpublished data),
- 10 Figure 4 Summary of protein classes categorized the trapped proteins using PANTHER algorism
- 11 77 trapped genes were successfully categorized at least one of 21 protein classes by using PANTHER
- gene ontology algorism. The details of the trapped genes classified in each protein class are listed in
- 13 Table 2.
- 14 Figure 5 Disease-causing human orthologues of the trapped genes involved in human genetic
- 15 disorders in multi-organ systems.
- Supplemental Table 1 Generation of 11 hundred independent lines by all three reading frames of
- 17 mRFP reporter in both RP2 and RP8 cassettes.
- 18 Supplemental Table 2 Disease-causing human orthologues of the trapped protein
- 19 Supplemental Figure 1 Representative expression patterns of mRFP fusion protein by RP2 and
- 20 RP8 integration

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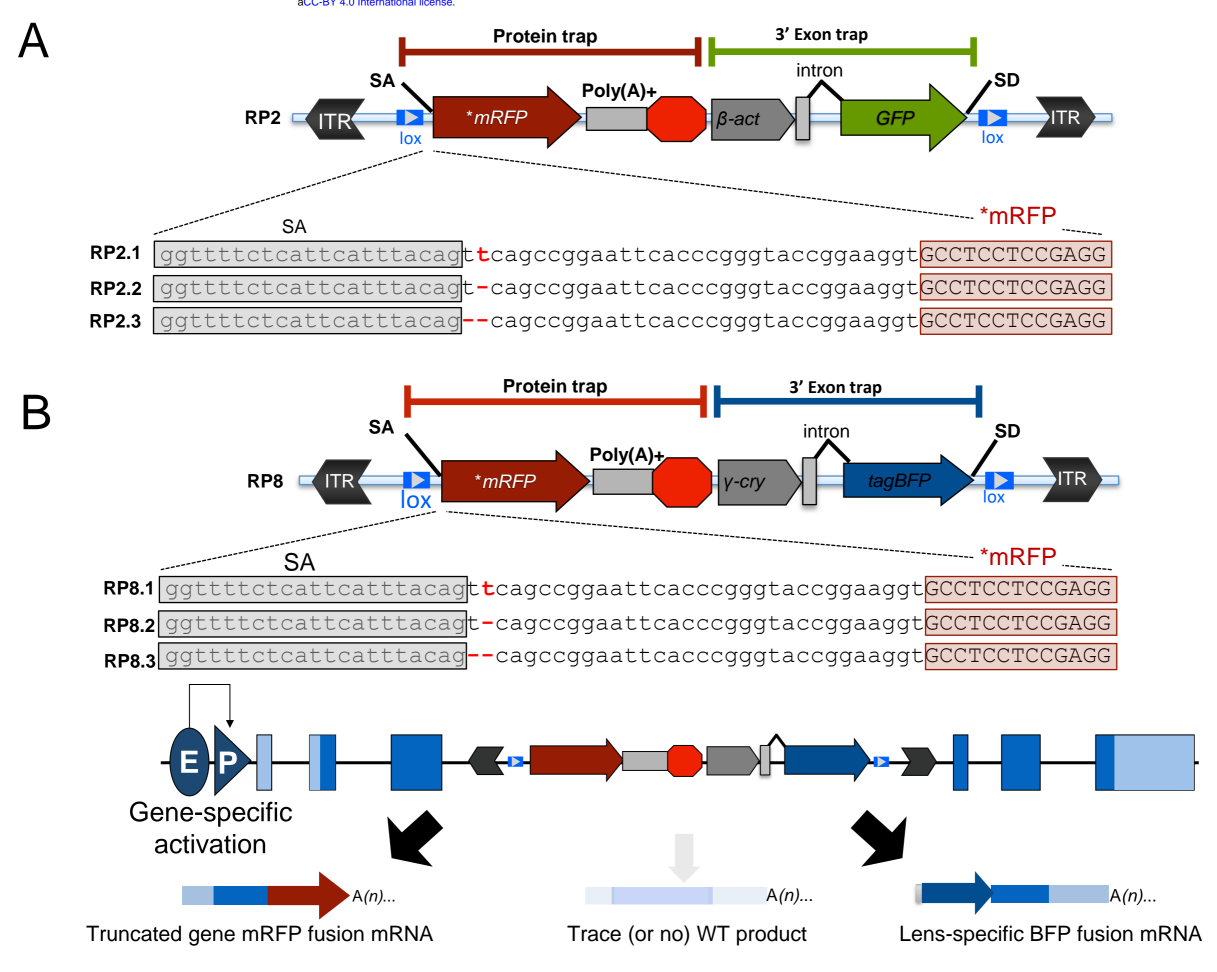


Fig. 1

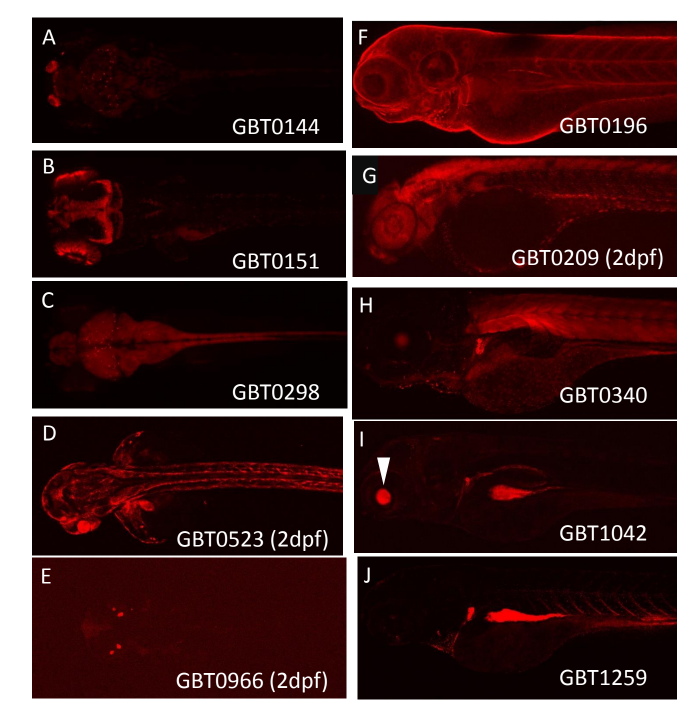


Fig.2

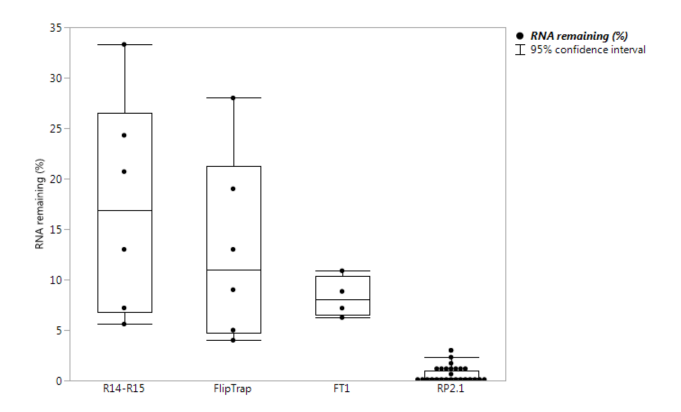


Fig. 3

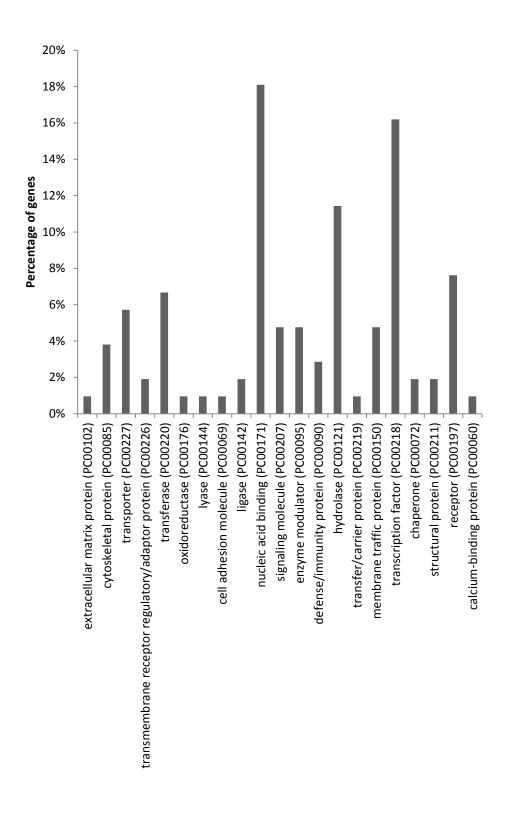


Figure 4

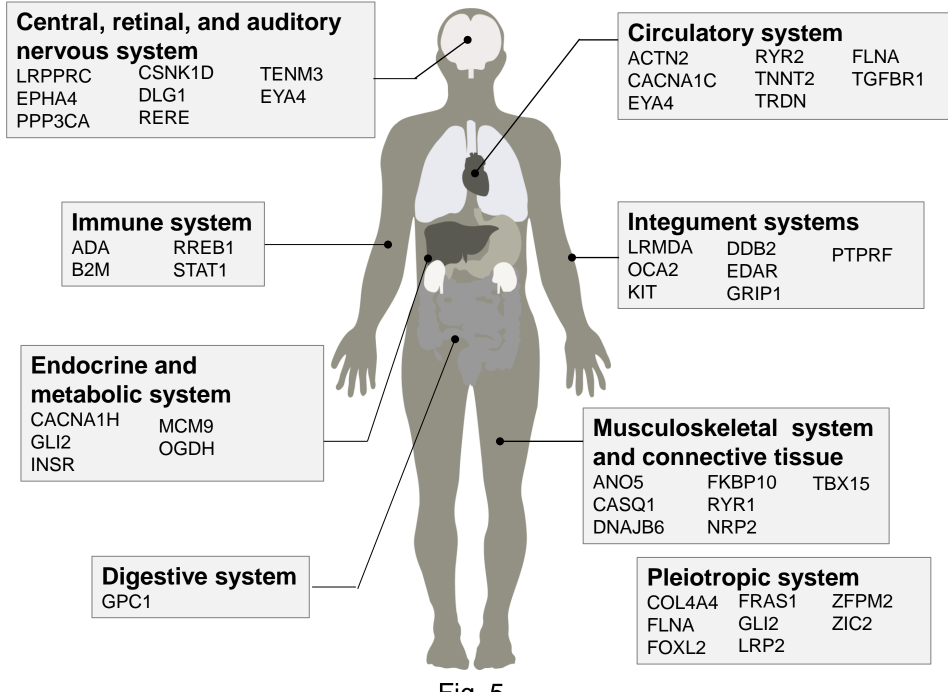


Fig. 5

	The number of genes		
Expression data exist	2 dpf	4 dpf	
zfishbook and ZFIN	45	10	
zfishbook	75	155	
ZFIN	20	2	
Not available	36	9	

Table 1

March Continue C	Protein Class	Mapped ID	Gene Symbol	Gene Name
Decide			•	
Dec Dec			gar1	H/ACA ribonucleoprotein complex subunit 1
200 GRIV-G001-04-04 rights 200 GRIV-G001-04-04 200 GRIV-G001-04-04-04-04-04-04-04-04-04-04-04-04-04-		ZDB-GENE-060503-160	•	
Page		ZDB-GENE-000405-1	pbx1a	Pbx1a homeodomain protein
Folia Micro Micro Folia 198 Page Folia Micro Micro Folia Micro Folia Micro Folia Micro Folia Micro Folia Micro Folia Micro Micro Folia		ZDB-GENE-060130-4	zfpm2a	Zinc finger protein, FOG family member 2a
DNA damage brinding protein 2 DNA GENE 20052-9-1 most mos		ZDB-GENE-050417-327	eef1a1b	Elongation factor 1-alpha
March Marc		ZDB-GENE-081104-328	enox1	Ecto-NOX disulfide-thiol exchanger 1
208.6FN-200329-1 thst5		ZDB-GENE-050419-169	ddb2	DNA damage-binding protein 2
2006 ENN-070314-013 1017 10401		ZDB-GENE-980526-306	msxc	Homeobox protein MSH-C
2006_CRN-0.0913-1.513			tbx15	
200 Series 200913-135 3				·
Description				
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2D8_GENE_06003-11 abcf1				
Dept. Sept. Sept				·
Description factor (PCOD219) Description factor (PCOD219)				
Post-Process Post				•
Properties Pro				
Vision Control (PCO0218) CDB-GENE-050913-92 CDB-GENE-050913-139 CDB-GENE-050913-139 CDB-GENE-050913-139 CDB-GENE-050913-149 CDB-GENE-050913-149 CDB-GENE-050913-146 CDB-GENE-050913-146 CDB-GENE-050913-146 CDB-GENE-050913-146 CDB-GENE-050913-146 CDB-GENE-050913-146 CDB-GENE-050913-142 CDB-GENE-050913-142 CDB-GENE-050913-133 CDB-GENE-050913-133 CDB-GENE-050913-133 CDB-GENE-050913-133 CDB-GENE-050913-133 CDB-GENE-050913-134 CDB-GENE-050913-134 CDB-GENE-050913-135 CDB-GENE-050913-136 CDB-GENE-05091	Rxiv preprint doi: https://doi.org	g/10.1101/630236; this version pos	sted May 7, 2019. The co	opyright holder for this preprint (which was not
ZDB-GENE-050413-261				proprint in perposality. It restricted available under
2DB-GENE-000015-1 pbx1a		•	wtip	Wilms tumor protein 1-interacting protein homolog
2DB-GENE-00010-4			•	
ZDB-GENE-060130-4				
ZDB-GENE-05019-146		ZDB-GENE-060130-4	•	·
DB-GENE-09013-1-66 bhlhe41		ZDB-GENE-000710-4	zic2a	ZIC family member 2 (Odd-paired homolog, Drosophila), A
ZDB-GENE-030131-6789 taffs		ZDB-GENE-061207-62	bcl11ba	B cell CLL/lymphoma 11Ba
Abmoebox protein MSH-C 2DB-GENE-020529-1		ZDB-GENE-050419-146	bhlhe41	BHLH protein DEC2
ZDB-GENE-020529-1 tp.		ZDB-GENE-030131-6789	taf6l	TAF6-like RNA polymerase II, p300/CBP-associated factor (PCAF)-associated factor
ZDB-GENE-070314-2		ZDB-GENE-980526-306	msxc	·
ZDB-GENE-030131-6317 dido1			tbx15	
Aphrolase PC00121				·
ZDB-GENE-980526-499 stat1a Castor zinc finger 1 Castor zinc zinc zinc zinc zinc zinc zinc zinc				
Aphrolase (PC00121) Table General Control of Contro				·
Aphrolase (PC00121) TOB-GENE-060512-241 TOB-GENE-040611-3 TOB-GENE-060503-150 TOB-GENE-040611-3 TOB-GENE-040611-3				
Aptrolase (PC00121)				
Adenosine deaminase, RNA-specific, B2 (non-functional) (Fragment)				
ZDB-GENE-040611-3	hvdrolase (PC00121)	ZDD GENE GHIIII HI	matesa	Tractical factor of activated 1 cells 3d
ZDB-GENE-060503-160 adarb2 Adenosine deaminase, RNA-specific, B2 (non-functional) (Fragment)	,	ZDB-GENE-040611-3	nrp2b	Neuropilin
ZDB-GENE-070112-1812 pnpla7a Patatin-like phospholipase domain-containing 7a ZDB-GENE-050417-327 eef1a1b Elongation factor 1-alpha ZDB-GENE-040718-393 ada Adenosine deaminase ZDB-GENE-140703-2 plpp2a Phospholipid phosphatase 2a ZDB-GENE-080818-1 ca16b Carbonic anhydrase XVI b ZDB-GENE-050517-31 abcf1 ATP-binding cassette, sub-family F (GCN20), member 1 ZDB-GENE-050503-530 ptprf Receptor-type tyrosine-protein phosphatase F ZDB-GENE-070117-757 parga Poly(ADP-ribose) glycohydrolase ENSDARG0000010758 capn12 Calpain 12 ZDB-GENE-041014-310 mcm9 DNA helicase MCM9 receptor (PC00197) ZDB-GENE-050419-65 v2rl1 Vomeronasal 2 receptor, I1 ZDB-GENE-050313-2427 col7a1 Collagen, type VII, alpha 1 ZDB-GENE-030131-2427 col7a1 Collagen, type VII, alpha 1 ZDB-GENE-030131-2427 tgfbr1b Serine/threonine-protein kinase receptor ZDB-GENE-091027-1 tgfbr1b Serine/threonine-protein kinase receptor ZDB-GENE-091027-1 tgfbr1b Serine/threonine-protein kinase receptor ZDB-GENE-030131-2049 mat2aa S-adenosylmethionine synthase ZDB-GENE-040426-1516 mboat7 Lysophospholipid acyltransferase 7 ZDB-GENE-040426-1516 mboat7 Lysophospholipid acyltransferase 7 ZDB-GENE-040403-11 kat2a Histone acetyltransferase KPZA ZDB-GENE-030131-205 csnk1da Casein kinase I isoform delta-A			•	·
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ENSDARG0000010758 capn12 Calpain 12 ZDB-GENE-041014-310 mcm9 DNA helicase MCM9 receptor (PC00197) ZDB-GENE-050419-65 v2rl1 Vomeronasal 2 receptor, l1 ZDB-GENE-060503-5 gabbr1b Gamma-aminobutyric acid (GABA) B receptor, 1b ZDB-GENE-070314-2 rxraa Retinoic acid receptor RXR-alpha-A ZDB-GENE-030131-2427 col7a1 Collagen, type VII, alpha 1 ZDB-GENE-03099-10 itgb1b Integrin beta ZDB-GENE-03099-10 tgb1b Serine/threonine-protein kinase receptor ZDB-GENE-091027-1 tgfbr1b Serine/threonine-protein kinase receptor ZDB-GENE-070905-3 crfb4 Cytokine receptor family member b4 transferase (PC00220) ZDB-GENE-030131-2049 mat2aa S-adenosylmethionine synthase ZDB-GENE-040426-1516 mboat7 Lysophospholipid acyltransferase 7 ZDB-GENE-121129-1 csgalnact1a Hexosyltransferase ZDB-GENE-060616-238 mgat5 Mannosyl (alpha-1,6-)-glycoprotein beta-1,6-N-acetyl-glucosaminyltransferase ZDB-GENE-080403-11 kat2a Histone acetyltransferase KAT2A ZDB-GENE-030131-825 csnk1da Casein kinase I isoform delta-A		ZDB-GENE-060503-530	ptprf	Receptor-type tyrosine-protein phosphatase F
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		ZDB-GENE-080403-11	_	
ZDB-GENE-091027-1 tgfbr1b Serine/threonine-protein kinase receptor		ZDB-GENE-030131-825	csnk1da	Casein kinase I isoform delta-A
-		ZDB-GENE-091027-1	tgfbr1b	Serine/threonine-protein kinase receptor

transporter (PC00227)			
	ZDB-GENE-130103-1	cacna1ha	Voltage-dependent T-type calcium channel subunit alpha
	ZDB-GENE-050419-65	v2rl1	Vomeronasal 2 receptor, l1
	ZDB-GENE-070705-417	ryr1b	Ryanodine receptor 1b (skeletal)
	ZDB-GENE-001127-2	atp1b2a	Sodium/potassium-transporting ATPase subunit beta
	ZDB-GENE-050517-31	abcf1	ATP-binding cassette, sub-family F (GCN20), member 1
	ZDB-GENE-070718-4	oca2	Oculocutaneous albinism II
on Turns of the design of the second		UCaZ	Oculocutarieous albinism ii
enzyme modulator (PC	-		Adamasina dagminasa DNA sasaifia D2 (sasa functional) (Functional)
	ZDB-GENE-060503-160	adarb2	Adenosine deaminase, RNA-specific, B2 (non-functional) (Fragment)
	ZDB-GENE-050417-327	eef1a1b	Elongation factor 1-alpha
	ZDB-GENE-030131-9805	ppp1r13ba	Protein phosphatase 1, regulatory subunit 13Ba
	ZDB-GENE-110411-39	si:dkey-222l13.1	Si:dkey-222l13.1
	ZDB-GENE-040426-1131	phactr4b	Phosphatase and actin regulator 4B
membrane traffic prot	ein (PC00150)		
	ZDB-GENE-030131-5290	napab	N-ethylmaleimide sensitive fusion protein attachment protein alpha
	ZDB-GENE-050522-134	syt5b	Synaptotagmin Vb
	ZDB-GENE-040718-281	vapal	VAMP (vesicle-associated membrane protein)-associated protein A,-like
	ZDB-GENE-050522-235	sncgb	Synuclein, gamma b (breast cancer-specific protein 1)
	ZDB-GENE-110411-39	si:dkey-222l13.1	Si:dkey-222l13.1
signaling molecule (DC			·
Rxiv preprint doi: https://doi.or	002077 g/10.1101/ <u>6</u> 30236; this yersion pos	sted May 7, 2019. The c	opyright holder for this preprint (which was not partianជាគ្រាស់ស្រាស់ ទីស្រាស់ ខេត្តដែលមិន្តាមហ៊ុន បាន
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	ZDB-GENE-080225-17.0 Into	-	Rho guanine nucleotide exchange factor (GEF) 25b
	ZDB-GENE-070615-10	nrg2a	Neuregulin 2a
	ZDB-GENE-041114-101	fgf13a	Fibroblast growth factor
	ZDB-GENE-040426-1793	fgf13b	Fibroblast growth factor
cytoskeletal protein (P	C00085)		
	ZDB-GENE-070112-1512	wasf3b	WAS protein family, member 3b
	ZDB-GENE-080215-23	map7d1b	MAP7 domain-containing 1b
	ZDB-GENE-060503-22	map7d1a	MAP7 domain-containing 1a
	ZDB-GENE-000626-1	tnnt2a	Troponin T type 2a (cardiac)
defense/immunity pro			· · · · · · · · · · · · · · · · · · ·
	ZDB-GENE-060503-160	adarb2	Adenosine deaminase, RNA-specific, B2 (non-functional) (Fragment)
	ZDB-GENE-040426-2136	b2ml	Beta-2-microglobulin
-l	ZDB-GENE-070905-3	crfb4	Cytokine receptor family member b4
chaperone (PC00072)			
	ZDB-GENE-040426-876	cct8	Chaperonin-containing TCP1, subunit 8 (theta)
	ZDB-GENE-050522-235	sncgb	Synuclein, gamma b (breast cancer-specific protein 1)
ligase (PC00142)			
	ZDB-GENE-090313-285	zmiz2	Zinc finger, MIZ-type-containing 2
	ZDB-GENE-070912-399	si:dkey-181m9.8	Si:dkey-181m9.8
structural protein (PC0	00211)		
	ZDB-GENE-030710-8	gpm6ab	Glycoprotein M6Ab (Fragment)
	ZDB-GENE-030710-9	gpm6ba	DMgamma1
transmembrane recen	tor regulatory/adaptor prot	0,	-
	ZDB-GENE-081030-10	cntn3a.1	Contactin 3a, tandem duplicate 1
	ZDB-GENE-031030-10 ZDB-GENE-010724-8	dlg1	Disks large homolog 1
cell adhesion molecule		αι δ±	NOW INTELLIGITION T
cen aunesion moiecule	•	itah1h	Integrin heta
autum adli dan er ertert	ZDB-GENE-030909-10	itgb1b	Integrin beta
extracellular matrix pr	•		Multiple FOE like demains Cl
-	ZDB-GENE-101112-3	megf6b	Multiple EGF-like-domains 6b
lyase (PC00144)		npr2	Guanylate cyclase
, ,	ZDB-GENE-141030-2	пртг	Guarry late eyelase
lyase (PC00144) oxidoreductase (PC001	176)	ΠριΖ	
, ,		enox1	Ecto-NOX disulfide-thiol exchanger 1
, ,	1 76) ZDB-GENE-081104-328	•	
oxidoreductase (PC001	1 76) ZDB-GENE-081104-328	•	
oxidoreductase (PC001	176) ZDB-GENE-081104-328 n (PC00219) ZDB-GENE-060628-2	enox1	Ecto-NOX disulfide-thiol exchanger 1
oxidoreductase (PC001 transfer/carrier protei	176) ZDB-GENE-081104-328 n (PC00219) ZDB-GENE-060628-2	enox1	Ecto-NOX disulfide-thiol exchanger 1

Table 2

	ZF NCBI		Human Orthologue	Human Orthologue
Line	gene ID	ZF gene symbol	NCBI Gene ID	Gene symbol
GBT0001	559053	casz1	54897	CASZ1
GBT0002	563408	sorbs2b	8470	SORBS2
GBT0005	570216	itgb1b	3688	ITGB1
GBT0007	794348		55051	NRDE2
GBT0010	30461	cdh11	1009	CDH11
GBT0016	58138	pbx1a	5087	PBX1
GBT0019	565629	kcnk10b	54207	KCNK10
GBT0020	569437	cntn3a.1	5067	CNTN3
GBT0021	571891	cntnap5b	129684	CNTNAP5
GBT0025	553277	dido1	11083	DIDO1
GBT0028	101882228	CABZ01057928.1	1729	DIAPH1
GBT0031	58071	tnnt2a	7139	TNNT2
GBT0033	100003333	Irch4	4034	LRCH4
GBT0034	567642	si:dkey-181m9.8		
GBT0035	559134	parga	8505	PARG
GBT0038	541513	srpx	8406	SRPX
GBT0039	559579	gabbr1,2	2550	GABBR1
GBT0040	58049	hoxa3a	3201	HOXA4
GBT0043	323266	cd99l2		CD99L2
GBT0046	64270	epha4b	2043	EPHA4
GBT0060	797527	crfb4	3588	IL10RB
GBT0067	559614	myom3	127294	МҮОМ3
GBT0070	555610	mcm9	254394	мсм9
GBT0073	406467	abcf1		ABCF1
GBT0077	406467	abcf1	23	ABCF1
GBT0078	558006	grip1	23426	
GBT0082	394037	cct8	10694	
GBT0091	100002220			ENOX1
GBT0094	403003	fgf13b	2258	FGF13
GBT0096		sh3glb2b		SH3GLB2
GBT0101	553277			DIDO1
GBT0103	100002190	cyth3a	9265	CYTH3
GBT0111	569081	rerea		RERE
GBT0113		cntn3a.1		CNTN3
GBT0125	100333685			CD302
GBT0126	405902	<u>'</u>		NRP2
GBT0131	100004133			DDB2
GBT0133	58077		7546	
GBT0135		bhlhe41		BHLHE41
GBT0137	100004503			EEF1A1
GBT0141		gpm6ba		GPM6B
GBT0142	100004850			AATK
GBT0143	566046	· ·	126374	
GBT0144	565405	unkl	64718	UNKL

GBT0145	560542	epn2	22905	EPN2
GBT0151	567446	nusap1	51203	NUSAP1
GBT0154	449761	si:ch211-163l21.8	57535	KIAA1324
GBT0156	563428	fras1	80144	FRAS1
GBT0157	724006	mgat5	4249	MGAT5
GBT0166	64269	atp1b2a	482	ATP1B2
GBT0168	492758	fgf13a	2258	FGF13
GBT0170	378997	glcci1a	113263	GLCCI1
GBT0172	337397	tmem30aa	55754	TMEM30A
GBT0175	569561	arhgef25b	115557	ARHGEF25
GBT0178	436919	ada	100	ADA
GBT0181	30155	tenm3	55714	TENM3
GBT0186	170581	cacna1c	775	CACNA1C
GBT0187	100007836	si:dkey-253d23.3		
GBT0189	562459		84330	ZNF414
GBT0190	100001699			BARHL2
GBT0195	562459		84330	ZNF414
GBT0196	559276			AHNAK
GBT0200	541428	znrd1	30834	ZNRD1
GBT0201		mhc1zca	3140	
GBT0202	569081	rerea		RERE
GBT0203	678534		150084	
GBT0204	492787			CADM4
GBT0205	100151756			FAM117A
GBT0208	641575			ACTN2
GBT0209	550559	'		DPH1
GBT0230	797651	*	83700	
GBT0231	266983			NEO1
GBT0232	553567	•	6861	
GBT0235	557195	* *		LRPPRC
GBT0237	100006951			NRG2
GBT0238		tgfbr1b		TGFBR1
GBT0239		map7d1b		MAP7D1
GBT0240	100093707			BCL11B
GBT0241	561900	•		FIP1L1
GBT0242	554131			TEX261
GBT0243		ppp1r13ba	23368	PPP1R13B
GBT0245	30526			
GBT0249	791742			B2M
GBT0250		ptprma		PTPRM
GBT0251	692279			FOXL2
GBT0255	114441			NCAM2
GBT0256		ppp3cb		PPP3CA
GBT0261		capn12		CAPN12
GBT0263	559150			HMCN1
GBT0268	100534737		203859	
GBT0270	568996	zfpm2a	23414	ZFPM2

Table 3_Human orthologues of GBT confirmed genes updated Apr8-2019.xlsx

CDT0274	F.C.F.4.7.2		F.F.700	NAAD7D4
GBT0271		map7d1a		MAP7D1
GBT0275	554270			COL4A4
GBT0281	797715	-		OGDH
GBT0283		sh3kbp1		SH3KBP1
GBT0286	797252	emid1	129080	EMID1
GBT0292	541428	znrd1	30834	ZNRD1
GBT0298	100007373	zgc:194659		
GBT0312	799867	ptprfb	5792	PTPRF
GBT0313	565460	csmd2	114784	CSMD2
GBT0316	324381	fkbp10b	60681	FKBP10
GBT0319	791987	ino80c	125476	INO80C
GBT0321	791987	ino80c	125476	INO80C
GBT0322		si:ch211-160j14.3		COL23A1
GBT0323	558149			ADARB2
GBT0325		megf6b		MEGF6
GBT0329		csgalnact1a		CSGALNACT1
GBT0329		nfatc3a		NFATC3
GBT0346	553730			LRMDA
GBT0348	570245			RYR1
GBT0345	569183	<u>'</u>		PTPRG
GBT0357		mat2aa		MAT2A
GBT0365	553679			SNCG
GBT0369		gpm6ab		GPM6A
GBT0309 GBT0380		znf1140	2023	GFIVIOA
			3791	KDD
GBT0383	554230			
GBT0389		mosmob		MOSMO
GBT0396	562557		56006	
GBT0397		phactr4b		PHACTR4
GBT0398		pnpla7a		PNPLA7
GBT0399		kirrel3a		KIRREL3
GBT0401	560003		23085	
GBT0402	566484			SCAF11
GBT0404		CABZ01045212.1	ENSGGOG00000007420	
GBT0409	100331745	-		NPR2
GBT0410	436819			VAPA
GBT0411		dnajb6b		DNAJB6
GBT0412	407710	· ·	23039	
GBT0415		arrdc1b		ARRDC1
GBT0416	322795	csrnp1b	64651	CSRNP1
GBT0419	555578	rxraa	6256	RXRA
GBT0422	245700	insrb	3643	INSR
GBT0424	566039	v2rl1		
GBT0425	550455	mrps18b	28973	MRPS18B
GBT0433	393599	ddb1	1642	DDB1
GBT0434	393950	gar1	54433	GAR1
GBT0435	558326	nrxn2a	9379	NRXN2
GBT0437	445226	casq1a	844	CASQ1
-			1	

Table 3_Human orthologues of GBT confirmed genes updated Apr8-2019.xlsx

GBT0503		stat1a		STAT1
GBT0505	572369	kirrel3b	84623	KIRREL3
GBT0510	553343	si:ch211-266g18.10	10345	TRDN
GBT0511	567419	oca2	4948	OCA2
GBT0513	65239	map2k6	5608	MAP2K6
GBT0520	553367		2817	GPC1
GBT0522	30708			FABP2
GBT0523		marcksl1a		MARCKSL1
GBT0525		mapk8ip1b		MAPK8IP1
_			10140	
GBT0527	386629			
GBT0528		zbtb16a		ZBTB16
GBT0534	327082			NAPA
GBT0545	563632			WASF3
GBT0552	562615	znf1015	163033	ZNF579
GBT0554	492500	gsto2	119391	GSTO2
GBT0570	30154	gli2a	2736	GLI2
GBT0572	568184	lrp2a	4036	LRP2
GBT0585	30256	kita	3815	KIT
GBT0591	100009635	dhx37	57647	DHX37
GBT0593	393716		6500	SKP1
GBT0599	768182	-	90550	
GBT0700	558875		64927	
GBT0700		magi2a		MAGI2
GBT0717 GBT0717	557363		10629	
GBT0717 GBT0722			22982	
	569986	<u> </u>		
GBT0726	100003611		55086	KADX
GBT0734		si:dkey-15h8.15		
GBT0750	557081			COL7A1
GBT0757	555517			KAT2A
GBT0760	110437953	_		ADGRL2
GBT0776	796370	edar	10913	EDAR
GBT0785	393509	mboat7	79143	MBOAT7
GBT0795	562459	znf414	84330	ZNF414
GBT0906	393638	timm50	92609	TIMM50
GBT0926	790941	dele1	9812	DELE1
GBT0936	541552	eya4	2070	EYA4
GBT0941	100333571	·	83637	ZMIZ2
GBT0951		macrod2		MACROD2
GBT0959	562529			FLNA
GBT0955	563806			PLPP2
GBT0900 GBT0972	114446	<u> </u>		DLG1
				ZFPM2
GBT0978		zfpm2a		
GBT0980	559150			HMCN1
GBT0993	557073	·	29965	
GBT1023		csnk1da		CSNK1D
GBT1027	557315			KIRREL2
GBT1033	555701	rreb1b	6239	RREB1

Table 3_Human orthologues of GBT confirmed genes updated Apr8-2019.xlsx

GBT1042	562146	pard3bb	117583	PARD3B
GBT1093	100126126	ryr2a	6262	RYR2
GBT1105	246222	tbx15	6913	TBX15
GBT1129	560875	cacna1ha	8912	CACNA1H
GBT1248	100148041	tnk2a	10188	TNK2
GBT1259	100537272	LOC100537272		
GBT1278	735249	selenos	55829	SELENOS
GBT1300	386856	pdgfra	5156	PDGFRA

Table 3

	Tagged Gene	Human Orthologue	Human Disease	Disease Models (Number of models)	Ref
ODTOG C	mb.:4:	DDV4	CONGENITAL ANOMALIES OF KIDNEY AND URINARY TRACT SYNDROME WITH ORWITHOUT HEARING LOSS	maus - /1)	
GBT0016	pbx1a	PBX1	ABNORMAL EARS OR DEVELOPMENTAL DELAY	mouse (1)	12
					18
					27
CDT0024	t	TAIAITO	CARDIONAVORATUV DU ATER 1D		17.
GBT0031	tnnt2a	TNNT2	CARDIOMYOPATHY DILATED 1D	mouse (5)	18
					16
					97
					11
					18
					10
					96
GBT0031	tnnt2a	TNNT2	CARDIOMYOPATHY FAMILIAL HYPERTROPHIC 2	mouse (9)	23
					14
GBT0078	grip1	GRIP1	FRASER SYNDROME 3	mouse (2)	16
					14
GBT0131	ddb2	DDB2	XERODERMA PIGMENTOSUM COMPLEMENTATION GROUP E	mouse (4)	15
					27
					18
GBT0133	zic?a	ZIC2	HOLOPROSENCEPHALY 5	mouse (3)	10
	bhlhe41	BHLHE41	SHORT SLEEPER		19
3010133	billile41	DUTUE41	SHORT SLEEPER	mouse (1)	_
					12
					12
					24
	1.				15
GBT0156	fras1	FRAS1	FRASER SYNDROME 1	mouse (5)	15
			SEVERE COMBINED IMMUNODEFICIENCY AUTOSOMAL RECESSIVE TCELL-NEGATIVE B CELL-NEGATIVE NK CELL-		
GBT0178	ada	ADA	NEGATIVE DUE TO ADENOSINEDEAMINASE DEFICIENCY	mouse (1)	94
GBT0186	cacna1c	CACNA1C	TIMOTHY SYNDROME	mouse (1)	21
GBT0240		BCL11B	IMMUNODEFICIENCY 49	zebrafish (1)	
				1	15
					14
GBT0251	foyl2a	FOXL2	Blepharophimosis ptosis and epicanthus inversus	mouse (3)	24
	ano5a	ANO5	MUSCULAR DYSTROPHY LIMB-GIRDLE AUTOSOMAL RECESSIVE 12	mouse (1)	26
JD10200	aliosa	ANUS	WIOSCOLAR DISTROPHI LIMB-GIRDLE AUTOSOMAL RECESSIVE 12	mouse (1)	
				(4)	10
GBT0270	· ·	ZFPM2	Tetralogy of Fallot	mouse (1)	12
GBT0270	zfpm2a	ZFPM2	DIAPHRAGMATIC HERNIA 3	mouse (1)	16
					21
	col4a4	COL4A4	ALPORT SYNDROME AUTOSOMAL RECESSIVE	manua (4)	1 2/
GBT0275	C014a4	COL4A4	ALPORT STINDROIVE AUTOSOWIAL RECESSIVE	mouse (4)	24
GBT0275	C014d4	COL4A4	ALPORT STINDROIVIE AUTOSOIVIAL RECESSIVE	mouse (4)	
GBT0275	C014a4	COL4A4	ALPORT STINDROIVIE AUTOSOIVIAL RECESSIVE	mouse (4)	25
				mouse (4)	25 19
				mouse (3)	25 19 7
			26FBNTRALLSAGFERSERSERSERSERSERSERSERSERSERSERSERSERSE		25 19 7: 27
ብዱፐ ዐ፭:4_{Åtp} ያቴ የህ ያሣሪ ^{is}	F.N.64.brg/10.1101/63 the author/funder, wh	802 BY R14s version posted May 7, no becomes the property of t	26FBNTRAUSAREIDISEASEIREFIMHA (WEIch was not 이 대한 사기에 마이어 한 이 아이어 한 아이어 이 아이어 이 아이어 아이어 아이어 아이어 아이어 아이어 아이어	mouse (3) mouse (1)	25 19 7! 27 23
මුසුT0348 _{tp} පුසු <i>T</i> 0396 ^{is} GBT0409	F.W.d.l.brg/10.1101/63 the author/funder, wh npr2	802 Bo RMs version posted May 7, 10 per granted bioRxiv a license to SV 200 By 4.0 International lid	ဥင်ဒေ့NTRAမ် မှာရှာနှေးမျှန်နေရှိမြောမြန်နေမှ was not ငှုံးမြောင်းမျှန်ရှာဖြောင်းမျှော်လည်းမှာဝန်းကြာချစ်ကျသို့ခဲ့ကြားဖြောက်ခြေ Acromesomelic dysplasia Maroteaux type	mouse (3) mouse (1) mouse (2)	25 19 7! 27 23 17
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ብሔፐ 03:4 _{Å tp} 336 ቸዊ 1596 is GBT 0409 GBT 0411	F.W.d.l.brg/10.1101/63 the author/funder, wh npr2	802 Bo RMs version posted May 7, 10 per granted bioRxiv a license to SV 200 By 4.0 International lid	ဥင်ဒေ့NTRAမ် မှာရှာနှေးမျှန်နေရှိမြောမြန်နေမှ was not ငှုံးမြောင်းမျှန်ရှာဖြောင်းမျှော်လည်းမှာဝန်းကြာချစ်ကျသို့ခဲ့ကြားဖြောက်ခြေ Acromesomelic dysplasia Maroteaux type	mouse (3) mouse (1) mouse (2)	25 19 7: 27 23 17 26
GBT0348tp GBT0409 GBT0411 GBT0422	the author/funder, when the author/funder, when the same	NPR2 DNAJB6 INSR	26FENTRAL CASE REPORTACION STANDERONE Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT)	25 19 7 27 23 17 26
GBT0449 GBT0409 GBT0422 GBT0511	npr2 dnajb6b insrb	NPR2 DNAJB6 INSR	26ENTRALSOBE DISEASE PROBLEM WAS not DEPAY PARISON OF THE PROBLEM	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1)	25 19 7: 27 23 17 26 Ma
GBT0449 GBT0409 GBT0422 GBT0511	the author/funder, when the author/funder, when the same	NPR2 DNAJB6 INSR	26FENTRAL CASE REPORTACION STANDERONE Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT)	25 19 7 27 23 17 26 Ma K.
GBT0409 GBT0411 GBT0422 GBT0511 GBT0572	npr2 dnajb6b insrb oca2 Irp2a	NPR2 DNAJB6 INSR OCA2 LRP2	26FINTRAL GABE IDISEAS FIRE MARCHEICH was not CHIPMAY TARION STANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 7: 27 23 17 26 Ma K.
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572	npr2 dnajb6b insrb oca2 lrp2a	NPR2 DNAJB6 INSR OCA2 LRP2 KIT	26日 TRAL CARE DISEASE DE MARION SANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 7 27 23 17 26 Ma K. 20 24 21
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572	npr2 dnajb6b insrb oca2 Irp2a	NPR2 DNAJB6 INSR OCA2 LRP2	26FINTRAL GABE IDISEAS FIRE MARCHEICH was not CHIPMAY THE RIPPER AND BRIAN PORTUPOR MARCHEICH WAS NOT CHISE. Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 72 27 23 17 26 Ma K. 20 24 21
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572	npr2 dnajb6b insrb oca2 lrp2a	NPR2 DNAJB6 INSR OCA2 LRP2 KIT	26日 TRAL CARE DISEASE DE MARION SANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 7: 27 23 17 26 Ma K. 20 24 21 20
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572	npr2 dnajb6b insrb oca2 lrp2a	NPR2 DNAJB6 INSR OCA2 LRP2 KIT	26日 TRAL CARE DISEASE DE MARION SANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 Irp2a kita kita	NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT	26日 TRAL CARE DISEASE DE MARION SANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22 18
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 Irp2a kita kita	NPR2 DNAJB6 INSR OCA2 LRP2 KIT	26日 TRAL CARE DISEASE DE MARION SANDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22 18
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 Irp2a kita kita	NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22 18 12
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 lrp2a kita kita	NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT	Gastrointestinal stromal tumor	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22 18 12
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 lrp2a kita kita	NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT	Gastrointestinal stromal tumor	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 72 23 17 26 Ma K. 20 24 21 20 16 22 18 12
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0572 GBT0585 GBT0585 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 lrp2a kita kita kita magi2a	DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2	GENTRAL CORF POSE ASSING MARCHER was not depart Part Part Part Part Part Part Part P	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 12 25
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 lrp2a kita kita kita magi2a	NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT	Gastrointestinal stromal tumor	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 12 25 18
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0572 GBT0585 GBT0585 GBT0585 GBT0585	npr2 dnajb6b insrb oca2 lrp2a kita kita kita magi2a	DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2	GENTRAL CORF POSE ASSING MARCHER was not depart Part Part Part Part Part Part Part P	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 12 25 18 19 10
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0585 GBT0750	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a	DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1	GENTRAL CORE IDISEASTINE MUSCULich was not SUPERIAL PROPERTIES AND STATE OF SEASTINE STATE OF SEASTIME STATE OF SEASTINE	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 10 17
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0585 GBT0750	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1	BOURTH Service of May 7, 100 Service of May 4.0 International life NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR	GENTRAL CORF IDISEAS IN CAMPORMATION SYNDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (4)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 10 17 9
GBT0348tp GBT0396 is GBT0409 GBT0411 GBT0422 GBT0572 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1	BOY BY Risks version posted May 7, 100 Sec granted bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR	GENTRAL CORE DISEASE OF MUSCULIC was not SHEAR T AND BRAIN WATPORMATION SYNDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (4) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 10 17 9 18
GBT0348tp GBT0396 is GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0785	npr2 dnajb6b insrb oca2 Irp2a kita kita magi2a col7a1 edar edar mboat7	BOY BY Rais version posted May 7, 100 Strong anted bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1) mouse (1) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23
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GBT0348tp SBT0396 GBT0409 GBT0411 GBT0422 GBT0511 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT07785 GBT0959	npr2 dnajb6b insrb oca2 Irp2a kita kita magi2a col7a1 edar edar mboat7	BOY BY Rais version posted May 7, 100 Strong anted bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1) mouse (1) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 10 17 9 18 23 16 15
GBT0348tp GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0785	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	24 25 19 7! 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9! 18 23 16 23 25 27
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GBT0348tp GBT0396 GBT0409 GBT0411 GBT0422 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT0785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24
GBT0348tp GBT0396 GBT0409 GBT0411 GBT0422 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT0785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24 23
GBT0348tp SBT0396 S GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT0776 GBT07785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24 23 22
GBT0348tp SBT0396 GBT0409 GBT0411 GBT0422 GBT0511 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT07785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 10 17 9 18 23 16 27 25 26 24 23 22 20
GBT0348tp SBT0396 GBT0409 GBT0411 GBT0422 GBT0511 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT07785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	SENTRAL CORE DISEASE OF MUSCLE: was not SHEAR T'S MY BRAINF WATER ON SYNDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1 ADVANCED SLEEP PHASE SYNDROME FAMILIAL 2	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24 23 22 20 18
GBT0348tp GBT0396 GBT0409 GBT0411 GBT0422 GBT0585 GBT0585 GBT0585 GBT0750 GBT0776 GBT0776 GBT0776 GBT0785 GBT0959	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24 23 22 20 18 16
GBT0348tp SBT0396 is GBT0409 GBT0411 GBT0422 GBT0511 GBT0572 GBT0585 GBT0585 GBT0750 GBT0776	npr2 dnajb6b insrb oca2 lrp2a kita kita magi2a col7a1 edar edar mboat7 flna	BOY BY Ris version posted May 7, 100 Separated bioRxiv a license to NPR2 DNAJB6 INSR OCA2 LRP2 KIT KIT MAGI2 COL7A1 EDAR EDAR EDAR MBOAT7 FLNA	SENTRAL CORE DISEASE OF MUSCLE: was not SHEAR T'S MY BRAINF WATER ON SYNDROME Acromesomelic dysplasia Maroteaux type autosomal dominant limb-girdle muscular dystrophy type 1 DONOHUE SYNDROME ALBINISM OCULOCUTANEOUS TYPE II DONNAI-BARROW SYNDROME MASTOCYTOSIS CUTANEOUS Piebald trait Gastrointestinal stromal tumor NEPHROTIC SYNDROME TYPE 15 recessive dystrophic epidermolysis bullosa hypohidrotic ectodermal dysplasia HAIR MORPHOLOGY 1 MENTAL RETARDATION AUTOSOMAL RECESSIVE 57 Periventricular nodular heterotopia 1 ADVANCED SLEEP PHASE SYNDROME FAMILIAL 2	mouse (3) mouse (1) mouse (2) zebrafish (1-NOT), mouse (1) mouse (2-NOT) rabbit (1) mouse (1) mouse (5) mouse (1) mouse (1)	25 19 75 27 23 17 26 Ma K. 20 24 21 20 16 22 18 19 10 17 9 18 23 16 15 27 25 26 24 23 22 20 18

Table 4

Line	Vector	Genome location	Genome version
GBT0115	RP2	chr8:12923125-12923133	ZV9
GBT0129	RP2	chr12:45451589-45451597	ZV9
GBT0148	RP2	chr16:55188302-55188310	ZV9
GBT0264	RP2	chr5:49960612-49960620	ZV9
GBT0506	RP2	chr18:46186350-46186358	ZV9
GBT0573	RP2	chr6:33630487-33630495	GRCz10
GBT0586	RP2	chr13:9112806-9112814	GRCz10
GBT0702	RP2	chr13:9112806-9112814	GRCz10
GBT0724	RP8	chr7:7633071-7633079	GRCz10
GBT0994	RP2	chr3:39627361-39627369	GRCz10
GBT1024	RP2	chr25:5175242-5175250	GRCz10
GBT1071	RP8	chr9:21818261-21818269	GRCz10
GBT1100	RP2	chr5:25509243-25509251	GRCz10
GBT1116	RP2	chr14:14724098-14724106	GRCz10
GBT1168	RP2	chr5:39664405-39664412	GRCz10

Table 5