# Supplementary Information

# Third-generation in situ hybridization chain reaction: multiplexed, quantitative, sensitive, versatile, robust

Harry M.T. Choi,<sup>1</sup> Maayan Schwarzkopf,<sup>1</sup> Mark E. Fornace,<sup>2</sup> Aneesh Acharya,<sup>1</sup> Georgios Artavanis,<sup>1</sup> Johannes Stegmaier,<sup>3,4,5</sup> Alexandre Cunha,<sup>3,6</sup> and Niles A. Pierce<sup>1,7,8,\*</sup>

# Contents

<b>S1</b>	Additional	materials and methods	6
	S1.1 Probe	set and amplifier details	6
	S1.2 Confe	ocal microscope settings	8
	S1.3 Flow	cytometer settings	9
	S1.4 Image	e analysis	11
	S1.4.1	Raw pixel intensities	11
	S1.4.2	Measurement of signal, background, and signal-to-background	11
	S1.4.3	Measurement of background components	12
	S1.4.4	Measurement of split-initiator HCR suppression	12
	S1.4.5	Normalized voxel intensities for qHCR imaging: analog mRNA relative quantitation with	
		subcellular resolution in an anatomical context	14
	S1.4.6	Dot detection and colocalization for dHCR imaging: digital mRNA absolute quantitation in	
		an anatomical context	14
	S1.5 Flow	cytometry data analysis	17
	S1.5.1	Raw cell intensities	17
	S1.5.2	Measurement of signal, background, signal-to-background, background components, and	
		split-initiator HCR suppression for transgenic targets	17
	S1.5.3	Measurement of signal, background, signal-to-background, background components, and	
		split-initiator HCR suppression for endogenous targets	17
	S1.5.4	Normalized single-cell intensities for qHCR flow cytometry: analog mRNA relative quanti-	
		tation for high-throughput expression profiling of human and bacterial cells	18
<b>S2</b>	Protocols fo	or in situ HCR v3.0	19
~ _	S2.1 Proto	cols for whole-mount chicken embryos	19
	S2.1.1	Preparation of fixed whole-mount chicken embryos	19
	S2.1.2	Buffer recipes for sample preparation	20
	S2.1.3	Multiplexed in situ HCR v3.0 using split-initiator probes	21
	S2.1.4	Buffer recipes for in situ HCR v3.0	22
	S2.1.5	Sample mounting for microscopy	23
	S2.1.6	Reagents and supplies	24

<sup>1</sup>Division of Biology & Biological Engineering, California Institute of Technology, Pasadena, CA 91125, USA. <sup>2</sup>Division of Chemistry & Chemical Engineering, California Institute of Technology, Pasadena, CA 91125, USA. <sup>3</sup>Center for Advanced Methods in Biological Image Analysis, Beckman Institute, California Institute of Technology, Pasadena, CA 91125, USA. <sup>4</sup>Institute for Automation & Applied Informatics, Karlsruhe Institute of Technology, Karlsruhe, Germany. <sup>5</sup>Institute of Imaging & Computer Vision, RWTH Aachen University, Aachen, Germany. <sup>6</sup>Center for Data-Driven Discovery, California Institute of Technology, Pasadena, CA 91125, USA. <sup>7</sup>Division of Engineering & Applied Science, California Institute of Technology, Pasadena, CA 91125, USA. <sup>8</sup>Weatherall Institute of Molecular Medicine, University of Oxford, Oxford OX3 9DS, UK. \*Email: niles@caltech.edu

	S2.2 Protocols for mammalian cells on a chambered slide	25
	S2.2.1 Preparation of fixed mammalian cells on a chambered slide	25
	S2.2.2 Buffer recipes for sample preparation	26
	S2.2.3 Multiplexed in situ HCR v3.0 using split-initiator probes	27
	S2.2.4 Buffer recipes for in situ HCR v3.0	28
	S2.2.5 Reagents and supplies	29
	S2.3 Protocols for mammalian cells in suspension	30
	S2.3.1 Preparation of fixed mammalian cells in suspension	30
	S2.3.2 Buffer recipes for sample preparation	31
	S2.3.3 Multiplexed in situ HCR v3.0 using split-initiator probes	32
	S2.3.4 Buffer recipes for in situ HCR v3.0	34
	S2.3.5 Reagents and supplies	35
	S2.4 Protocols for bacteria in suspension	36
	S2.4.1 Preparation of fixed bacteria in suspension	36
	S2.4.2 Buffer recipes for sample preparation	37
	S2.4.3 Multiplexed in situ HCR v3.0 using split-initiator probes	38
	S2.4.4 Buffer recipes for in situ HCR v3.0	40
	S2.4.5 Reagents and supplies	41
<b>S</b> 3	Additional studies	42
	S3.1 Validation of split-initiator HCR suppression in vitro and in situ (cf. Figure 2)	42
	S3.2 In situ validation of automatic background suppression with split-initiator probes in whole-mount	
	chicken embryos (cf. Figure 3)	44
	S3.2.1 Measurement of background and signal-to-background for unoptimized standard and split-	
	initiator probe sets as a function of probe set size	45
	S3.2.2 Measurement of background and signal-to-background for standard and split-initiator probes	
	with identical target-binding domains	49
	S3.2.3 Measurement of signal, background, signal-to-background, background components, and	
	split-initiator HCR suppression	51
	S3.3 Multiplexed 4-channel mRNA imaging with high signal-to-background in whole-mount chicken	50
	embryos (cf. Figure 4)	53
	S3.4 qHCR imaging: analog mRNA relative quantitation with subcellular resolution in whole-mount	
		55
	S3.4.1 Testing for a crowding effect $\dots \dots \dots$	33
	S3.4.2 Redundant 2-channel detection of $Dmbx1$	60 61
	S3.4.5 Redundant 2-channel detection of <i>EprA4</i>	01
	ss.s In situ vandation of automatic background suppression with spin-initiator probes for mkinA now	62
	S2.5.1 Massurement of signal background signal to background background components and	05
	solution in the submeasurement of signal, background, signal-to-background, background components, and solution in the submeasurement of signal for $d2aCEP$ transgenic target in HEK calls	63
	S3.5.2 Measurement of signal background signal to background background components and	05
	split initiator HCP suppression for <i>CAPDH</i> and genous target in HEK cells	65
	Solution of signal background signal to background background components and	05
	split-initiator HCR suppression for eGEP transgenic target in E coli	67
	S3.6 aHCR flow cytometry: analog mRNA relative quantitation for high-throughput expression profiling	07
	of human and hacterial cells (cf. Figure 6)	69
	S3 6.1 Testing for a crowding effect	60
	S3.6.2 Redundant 2-channel detection of <i>GAPDH</i> endogenous target in HFK cells	70
	S3.6.3 Redundant 2-channel detection of <i>PGK1</i> endogenous target in HEK cells	72
	S3.6.4 Redundant 2-channel detection of fusA endogenous target in F coli	7 <u>4</u>
		/ -r

	S	S3.6.5 Multiplexed 2-channel detection of <i>GAPDH</i> and <i>PGK1</i> endogenous targets in HEK cells 70							
	S	3.6.6 Multiplexed 2-channel detection of <i>fusA</i> and <i>icd</i> endogenous targets in <i>E. coli</i>	78						
	S3.7	4.7 dHCR imaging: digital mRNA absolute quantitation in an anatomical context (cf. Figure 7) 80							
	5	33.7.1 Redundant 2-channel detection of single <i>BRAF</i> mRNAs in HEK cells using in situ HCR v3.0	80						
	S	3.7.2 Redundant 2-channel detection of single <i>Dmbx1</i> mRNAs in whole-mount chicken embryos							
		using in situ HCR v3.0	81						
	S	33.7.3 Redundant 2-channel detection of single <i>kdrl</i> mRNAs in whole-mount zebrafish embryos							
		using in situ HCR v2.0 (Shah <i>et al.</i> , 2016) $\ldots$ $\ldots$ $\ldots$ $\ldots$ $\ldots$ $\ldots$ $\ldots$ $\ldots$	82						
<b>S4</b>	Prob	e sequences	83						
	S4.1	Standard probes for Figures 3, S5, and S7	84						
	S4.2	Split-initiator probes for Figures 3, S6, S8, S9, and S10	85						
	S4.3	Standard probes for Figures S9 and S10	86						
	S4.4	Split-initiator probes for Figure S11	88						
	S4.5	Split-initiator probes for Figures 4 and S12	89						
	S4.6	Split-initiator probes for Figures S13–S17	93						
	S4.7	Split-initiator probes Figures 5, S18, and S19	95						
	S4.8	Split-initiator probes for Figure 6, S20–S28	99						
	S4.9	Split-initiator probes for Figures 7, S29, and S30	108						

# List of Figures

<b>S</b> 1	Illustration of gates used for flow cytometry analysis of HEK cells	10
S2	Illustration of gates used for flow cytometry analysis of <i>E. coli</i>	10
<b>S</b> 3	Test tube validation of split-initiator HCR suppression for amplifiers B1, B2, B4, and B5 (cf. Figure 2)	42
<b>S</b> 4	Test tube validation of split-initiator HCR suppression for amplifier B3 (cf. Figure 2)	43
<b>S</b> 5	Measurement of background for unoptimized standard probe sets as a function of probe set size (cf.	
~~~	Figure 3A)	45
<b>S</b> 6	Measurement of background for unoptimized split-initiator probe sets as a function of probe set size	10
	(cf. Figure 3A)	46
<b>S</b> 7	Measurement of signal and background for unoptimized standard probe sets as a function of probe	
	set size (cf. Figure 3B)	47
<b>S</b> 8	Measurement of signal and background for unoptimized split-initiator probe sets as a function of	
	probe set size (cf. Figure 3B)	48
S9	Measurement of background for standard and split-initiator probes with identical target-binding do-	
	mains	49
S10	Measurement of signal and background for standard and split-initiator probes with identical target-	
	binding domains	50
S11	Measurement of signal and background, background components, and split-initiator HCR suppression	51
S12	Measurement of signal and background for multiplexed 4-channel mRNA imaging (cf. Figure 4)	53
S13	Comparison of signal intensity distributions for individual and simultaneous imaging of <i>EphA4</i> and	
	<i>Egr2</i>	55
S14	Characterizing signal plus background for <i>EphA4</i> and <i>Egr2</i> in a 2-target experiment	56
S15	Characterizing signal plus background for <i>EphA4</i> in a 1-target experiment	57
S16	Characterizing signal plus background for <i>Egr2</i> in a 1-target experiment	58
S17	Characterizing background for <i>EphA4</i> and <i>Egr2</i>	59
S18	Redundant 2-channel detection of <i>Dmbx1</i>	60
S19	Redundant 2-channel detection of <i>EphA4</i> (cf. Figure 5)	61
S20	Measurement of signal and background, background components, and split-initiator HCR suppres-	
	sion for <i>d2eGFP</i> transgenic target in HEK cells (cf. Figure 6A)	63

S21	Measurement of signal and background, background components, and split-initiator HCR suppression for <i>GAPDH</i> endogenous target in HEK cells	65
S22	Measurement of signal and background, background components, and split-initiator HCR suppression for <i>eGFP</i> transgenic target in <i>E. coli</i> (cf. Figure 6A)	67
S23	Comparison of signal intensity distributions for individual and multiplexed floHCR of <i>GAPDH</i> and <i>ACTB</i>	69
S24	Measurement of signal and background, background components, and split-initiator HCR suppres-	-0
S25	sion for redundant 2-channel detection of <i>GAPDH</i> endogenous target (cf. Figure 6B)	70
	sion for redundant 2-channel detection of <i>PGK1</i> endogenous target	72
S26	Measurement of signal and background, background components, and split-initiator HCR suppression for redundant 2 channel detection of <i>fusA</i> endogenous target (of Figure 6R)	74
S27	Measurement of signal and background, background components, and split-initiator HCR suppres-	/+
<b>GQ</b> 0	sion for multiplexed 2-channel detection of <i>GAPDH</i> and <i>PGK1</i> endogenous targets	76
\$28	sion for multiplexed 2-channel detection of <i>fusA</i> and <i>icd</i> endogenous targets	78
S29	Redundant 2-channel detection of single <i>BRAF</i> mRNAs in HEK cells using in situ HCR v3.0 (cf.	
<b>GQQ</b>	Figure 7A)	80
\$30	Redundant 2-channel detection of single <i>Dmbx1</i> mRNAs in whole-mount chicken embryos using in situ HCR v3.0 (cf. Figure 7B)	81
S31	Analysis of single-molecule mRNA imaging in whole-mount zebrafish embryos using in situ HCR	00
	v2.0	82

# List of Tables

C 1	Secure and for selectedies (of Eigen 2)	6
51	Sequences for get studies (cf. Figure 2)	0
<b>S</b> 2	Organisms, target mRNAs, probe sets, amplifiers, and figure numbers for in situ HCR experiments.	/
<b>S</b> 3	Confocal microscope settings	8
<b>S</b> 4	Flow cytometer settings	9
S5	Experiment types for qHCR imaging using in situ HCR v3.0	12
<b>S</b> 6	Parameters used for dot detection in dHCR images.	16
<b>S</b> 7	Experiment types for flow cytometry using in situ HCR v3.0 with a transgenic target mRNA	17
<b>S</b> 8	Experiment types for flow cytometry using in situ HCR v3.0 with an endogenous target mRNA	18
S9	In situ validation of split-initiator HCR suppression	42
S10	Estimated background for standard and split-initiator probes sets as a function of probe set size (cf.	
	Figure 3A)	46
<b>S</b> 11	Estimated signal-to-background for standard and split-initiator probes sets as a function of probe set	
	size (cf. Figure 3B)	48
S12	Estimated background for standard and split-initiator probes with identical target-binding domains .	49
S13	Estimated signal-to-background for standard and split-initiator probes with identical target-binding	
	domains	50
S14	Estimated signal-to-background, background components, and split-initiator HCR suppression	52
S15	Estimated signal-to-background for multiplexed 4-channel mRNA imaging	54
S16	Estimated signal-to-background for redundant 2-channel detection of <i>Dmbx1</i> and <i>EphA4</i>	62
S17	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	d2eGFP transgenic target in HEK cells (cf. Figure 6A)	64
S18	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	GAPDH endogenous target in HEK cells	66
S19	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	<i>eGFP</i> transgenic target in <i>E. coli</i> (cf. Figure 6A)	68

S20	Estimated signal-to-background, background components, and split-initiator HCR suppression for redundant 2-channel detection of <i>GAPDH</i> endogenous target (cf. Figure 6B)	71
S21	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	redundant 2-channel detection of <i>PGK1</i> endogenous target	73
S22	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	redundant 2-channel detection of <i>fusA</i> endogenous target (cf. Figure 6B)	75
S23	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	multiplexed 2-channel detection of GAPDH and PGK1 endogenous targets	77
S24	Estimated signal-to-background, background components, and split-initiator HCR suppression for	
	multiplexed 2-channel detection of <i>fusA</i> and <i>icd</i> endogenous targets	79
S25	Dot colocalization fractions for redundant 2-channel detection of single BRAF mRNAs in HEK cells	
	using in situ HCR v3.0 (cf. Figure 7A)	80
S26	Dot colocalization fractions for redundant 2-channel detection of single Dmbx1 mRNAs in whole-	
	mount chicken embryos using in situ HCR v3.0 (cf. Figure 7B)	81
S27	Dot colocalization fractions for redundant 2-channel detection of single kdrl mRNAs in whole-	
	mount zebrafish embryos using in situ HCR v3.0	82

# S1 Additional materials and methods

HCR Amplifier	Oligo	Length (nt)	Sequence $(5' \text{ to } 3')$	Figures
B1-Alexa647	I1	36	gAggAgggCAgCAAACgggAAgAgTCTTCCTTTACg	S3
	P1	45	gAggAgggCAgCAAACgg <mark>AAAgACgTTgTggCTgTTgTAgTTgTA</mark>	S3
	P2	45	CgTTCTTCTgCTTgTCggCCATgATTAgAAgAgTCTTCCTTTACg	S3
B2-Alexa647	I1 P1 P2	36 45 45	36CCTCgTAAATCCTCATCAATCATCCAgTAAACCgCCS45CCTCgTAAATCCTCATCAAAAgACgTTgTggCTgTTgTAgTTgTAS45CgTTCTTCTgCTgCCgCCATgATAAATCATCCAgTAAACCgCCS	
B3-Alexa647	I1	36	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	2, S4
	P1	45	gTCCCTgCCTCTATATCT <b>TTAgACgTTgTggCTgTTgTAgTTgTA</b>	2, S4
	P2	45	CgTTCTTCTgCTTgTCggCCATgATTTCCACTCAACTTTAACCCg	2, S4
B4-Alexa647	I1	36	CCTCAACCTACCTCCAACTCTCACCATATTCgCTTC	S3
	P1	45	CCTCAACCTACCTCCAACAAAgACgTTgTggCTgTTgTAgTTgTA	S3
	P2	45	CgTTCTTCTgCTTgTCggCCATgATATTCTCACCATATTCgCTTC	S3
B5-Alexa647	I1	36	CTCACTCCCAATCTCTATCTACCCTACAAATCCAAT	\$3
	P1	45	CTCACTCCCAATCTCTATAAAgACgTTgTggCTgTTgTAgTTgTA	\$3
	P2	45	CgTTCTTCTgCTTgTCggCCATgATAACTACCCTACAAATCCAAT	\$3

# S1.1 Probe set and amplifier details

Table S1. Sequences for gel studies (cf. Figure 2). Initiator sequence in green, spacer sequence in blue, target-binding sequence in black. In all cases, the Target is: 5'-TACAACTACAACAGCCACAACGTCTATATCATGGCCGACAAGCAGAAGAACG-3'.

		Standard	Split-initiator			
		probes	probe pairs	HCR	Amplification	
Organism	Target	(v2.0)	(v3.0)	amplifier	time	Figures
G. gallus domesticus	Sox10	5, 10, 20		B3-Alexa647	overnight	3, \$5, \$7
	Sox10		5, 10, 20	B3-Alexa647	overnight	3, S6, S8
	Sox10		20	B3-Alexa647	overnight	S9, S10
	Sox10	20		B3-Alexa647	overnight	S9, S10
	EphA4		20	B2-Alexa647	overnight	S11
	FoxD3		12	B4-Alexa488	overnight	4, S12
	Dmbx1		20	B1-Alexa514	overnight	4, S12
	Sox10		20	B3-Alexa546	overnight	4, S12
	EphA4		20	B2-Alexa647	overnight	4, S12
	EphA4		20	B1-Alexa546	overnight	S13, S14, S15
	Egr2		20	B3-Alexa647	overnight	S13, S14, S16
	Dmbx1		20	B1-Alexa546	overnight	S18
	Dmbx1		20	B2-Alexa647	overnight	S18
	EphA4		20	B1-Alexa546	overnight	5, S19
	EphA4		20	B2-Alexa647	overnight	5, S19
H. sapiens sapiens	d2eGFP		12	B3-Alexa594	overnight	6A, S20
	GAPDH		10	B5-Alexa488	overnight	6B, S23, S24
	GAPDH		10	B4-Alexa594	overnight	6B, S1, S21, S24
	ACTB		10	B2-Alexa594	overnight	S23
	PGK1		18	B1-Alexa488	overnight	S25
	PGK1		18	B2-Alexa594	overnight	S25, S27
	GAPDH		10	B4-Alexa488	overnight	S27
E. coli	eGFP		12	B3-Alexa594	overnight	6A, S22
	fusA		18	B3-Alexa488	overnight	6B, S26, S28
	fusA		18	B2-Alexa594	overnight	6B, S2, S26
	icd		20	B1-Alexa594	overnight	S28
H. sapiens sapiens	BRAF		23	B3-Alexa647	45 min	7A, S29
	BRAF		23	B4-Alexa546	45 min	7A, S29
G. gallus domesticus	Dmbx1		25	B1-Alexa594	90 min	7B, S30
	Dmbx1		25	B2-Alexa647	90 min	7B, S30

Table S2. Organisms, target mRNAs, probe sets, amplifiers, and figure numbers for in situ HCR experiments.

# S1.2 Confocal microscope settings

Target	Fluorophore	Laser (nm)	Beam Splitter	Filter (nm)	Pixel size $(x \times y \times z \ \mu m)$	Voxel size $(x \times y \times z \ \mu m)$	Focal planes	Figures
EmbAd	Aleve647	()	MDS 499/561/622	650 690	$\frac{(\omega \times g \times \lambda \mu m)}{0.200 \times 0.200}$	( <i>w</i> / <i>y</i> / <i>z p</i> )	1	C11
EpnA4	Alexa047	(22	MDS 488/501/055	030-089	$0.208 \times 0.208 \times 2.7$		1	311
Sox10	Alexa64/	633	MBS 488/561/633	650–699	$0.415 \times 0.415 \times 2.7$		1	3, 85–810
FoxD3	Alexa488	488	MBS 488/561/633	491–525	$0.664 \times 0.664 \times 4$		1	4, S12
Dmbx1	Alexa514	514	MBS 458/514	546-564	$0.664 \times 0.664 \times 4$		1	4, S12
Sox10	Alexa546	561	MBS 488/561/633	573-612	$0.664 \times 0.664 \times 4$		1	4, S12
EphA4	Alexa647	633	MBS 488/561/633	654–687	$0.664 \times 0.664 \times 4$		1	4, S12
Dmbx1	Alexa546	561	MBS 488/561/633	563–592	$0.208 \times 0.208 \times 2.7$	$2.1 \times 2.1 \times 2.7$	1	5, S18
Dmbx1	Alexa647	633	MBS 488/561/633	650–689	$0.208 \times 0.208 \times 2.7$	$2.1\times2.1\times2.7$	1	5, S18
EphA4	Alexa546	561	MBS 488/561/633	563-592	$0.208 \times 0.208 \times 2.7$	$2.1 \times 2.1 \times 2.7$	1	5, S19
EphA4	Alexa647	633	MBS 488/561/633	650–689	$0.208 \times 0.208 \times 2.7$	$2.1\times2.1\times2.7$	1	5, S19
BRAF	Alexa546	561	MBS 405/488/561/640 (T10/R90)	566-623	$0.0624 \times 0.0624 \times 0.42$		17	7A, S29
BRAF	Alexa647	640	MBS 405/488/561/640 (T10/R90)	656-700	$0.0624 \times 0.0624 \times 0.42$		17	7A, S29
	DAPI	405	MBS 405/488/561/640 (T10/R90)	410-470	$0.0624 \times 0.0624 \times 0.42$		17	S29
Dmbx1	Alexa594	561	MBS 405/488/561/640 (T10/R90)	580-647	$0.099\times0.099\times0.420$		22	7B, S30
Dmbx1	Alexa647	640	MBS 405/488/561/640 (T10/R90)	645-700	$0.099 \times 0.099 \times 0.420$		22	7B, S30
EphA4	Alexa546	561	MBS 488/561/633	563-592	$0.415 \times 0.415 \times 2.7$	$2.1 \times 2.1 \times 2.7$	1	S14–S17
Egr2	Alexa647	633	MBS 488/561/633	650–689	$0.415 \times 0.415 \times 2.7$	$2.1\times2.1\times2.7$	1	S14–S17

Table S3. Confocal microscope settings.

# S1.3 Flow cytometer settings

		Laser		Emission	
Target	Fluorophore	(nm)	Filter	(nm)	Figures
d2eGFP	Alexa594	561	Y2	615/20	6A, S20
GAPDH	Alexa488	488	<b>B</b> 1	525/50	6B, S23, S24, S27
GAPDH	Alexa594	561	Y2	615/20	6B, S1, S21, S24
ACTB	Alexa594	561	Y2	615/20	S23
PGK1	Alexa488	488	B1	525/50	S25
PGK1	Alexa594	561	Y2	615/20	S25, S27
eGFP	Alexa594	561	Y2	615/20	6A, S22
fusA	Alexa488	488	<b>B</b> 1	525/50	6B, S26, S28
fusA	Alexa594	561	Y2	615/20	6B, S2, S26
icd	Alexa594	561	Y2	615/20	S28

Table S4. Flow cytometer settings.



**Figure S1. Illustration of gates used for flow cytometry analysis of HEK cells.** (A) Scatter plots for ungated data. Top: side scatter area (SSC-A) vs forward scatter area (FSC-A). Bottom: forward scatter height (FSC-H) vs forward scatter area (FSC-A). (B) Scatter plots after applying one gate. Top: gate on FSC-A vs SSC-A to remove debris and select cells. Bottom: gate on FSC-A vs FSC-H to remove clumps of cells and select single cells. (C) Scatter plots after applying both gates. (D) Signal plus background for ungated (top) and gated (bottom) samples. Target: *GAPDH*. Probe set: 10 split-initiator probe pairs. Amplifier: B4-Alexa594. The depicted gates were used for the SIG+BACK data in Figure S21.



**Figure S2. Illustration of gates used for flow cytometry analysis of** *E. coli*. (A) Scatter plots for ungated sample (top) and gated sample (bottom): side scatter area (SSC-A) vs forward scatter area (FSC-A). (B) Signal plus background for ungated sample (top) and gated sample (bottom). Target: *fusA*. Probe set: 18 split-initiator probe pairs. Amplifier: B2-Alexa594. The depicted gate was used for the SIG+BACK data in Figure S26.

# S1.4 Image analysis

We build on an image analysis framework developed over a series of publications (Choi *et al.*, 2010, 2014, 2016; Trivedi *et al.*, 2018). For convenience, here we provide a self-contained description of the details relevant to the present work.

# S1.4.1 Raw pixel intensities

The total fluorescence within a pixel is a combination of signal and background. Fluorescent background (BACK) arises from three sources in each channel:

- autofluorescence (AF): fluorescence inherent to the sample,
- non-specific detection (NSD): probes that bind non-specifically in the sample and subsequently trigger HCR amplification,
- non-specific amplification (NSA): HCR hairpins that bind non-specifically in the sample.

Fluorescent signal (SIG) in each channel corresponds to:

• signal (SIG): probes that hybridize specifically to the target mRNA and subsequently trigger HCR amplification.

For pixel i of replicate embryo n, we denote the background

$$X_{n,i}^{\text{BACK}} = X_{n,i}^{\text{NSD}} + X_{n,i}^{\text{NSA}} + X_{n,i}^{\text{AF}},$$
(S1)

the signal:

$$X_{n,i}^{\mathrm{SIG}},$$
 (S2)

and the total fluorescence (SIG+BACK):

$$X_{n,i}^{\text{SIG+BACK}} = X_{n,i}^{\text{SIG}} + X_{n,i}^{\text{BACK}}.$$
(S3)

## S1.4.2 Measurement of signal, background, and signal-to-background

For each target mRNA, background (BACK) is characterized for pixels in a representative rectangular region of no- or low-expression and the combination of signal plus background (SIG+BACK) is characterized for pixels in a representative rectangular region of high expression (e.g., Figures 3C, S7A, S8A, S10A, and S12A). For the pixels in these regions, we characterize the distribution by plotting an intensity histogram (e.g., Figures 3D, S7B, S8B, S10B, and S12B) and characterize average performance by calculating the mean pixel intensity ( $\bar{X}_n^{\text{BACK}}$  or  $\bar{X}_n^{\text{SIG+BACK}}$  for replicate embryo *n*). Performance across replicate embryos is characterized by calculating the sample means ( $\bar{X}^{\text{BACK}}$  and  $\bar{X}^{\text{SIG+BACK}}$ ) and standard errors ( $s_{\bar{X}BACK}$  and  $s_{\bar{X}SIG+BACK}$ ). The mean signal is then estimated as

$$\bar{X}^{\text{SIG}} = \bar{X}^{\text{SIG+BACK}} - \bar{X}^{\text{BACK}} \tag{S4}$$

with the standard error estimated via uncertainty propagation as

$$s_{\bar{X}}$$
SIG  $\leq \sqrt{(s_{\bar{X}}$ SIG+BACK)^2 + (s\_{\bar{X}}BACK)^2}. (S5)

The signal-to-background ratio is estimated as:

$$\bar{X}^{\text{SIG/BACK}} = \bar{X}^{\text{SIG}} / \bar{X}^{\text{BACK}} \tag{S6}$$

with standard error estimated via uncertainty propagation as

$$s^{\text{SIG/BACK}} \leq \bar{X}^{\text{SIG/BACK}} \sqrt{\left(\frac{s_{\bar{X}^{\text{SIG}}}}{\bar{X}^{\text{SIG}}}\right)^2 + \left(\frac{s_{\bar{X}^{\text{BACK}}}}{\bar{X}^{\text{BACK}}}\right)^2}.$$
(S7)

These upper bounds on estimated standard errors hold under the assumption that the correlation between SIG and BACK is non-negative. Tables S10–S16 display signal, background, and/or signal-to-background values to characterize the performance of in situ HCR v3.0 within whole-mount chicken embryos.

			Reagents		
	Experiment type	Quantity	Probes	Hairpins	Expression region
Α	1	SIG+NSD+NSA+AF = SIG+BACK	odd + even	$\checkmark$	high
	1	NSD+NSA+AF = BACK	odd + even	$\checkmark$	no/low
В	2	NSA+AF		$\checkmark$	high
	3	AF			high
С	4	$SIG^{odd}$ + $NSD^{odd}$ + $NSA$ + $AF$ = $SIG^{odd}$ + $BACK^{odd}$	odd	$\checkmark$	high
	4	$NSD^{odd} + NSA + AF = BACK^{odd}$	odd	$\checkmark$	no/low
	5	SIG <sup>even</sup> +NSD <sup>even</sup> +NSA+AF = SIG <sup>even</sup> +BACK <sup>even</sup>	even	$\checkmark$	high
	5	$NSD^{even}$ + $NSA$ + $AF$ = $BACK^{even}$	even	$\checkmark$	no/low

**Table S5. Experiment types for qHCR imaging using in situ HCR v3.0.** (A) Characterize signal, background, and signal-tobackground. (B) Characterize components of background (AF, NSA, NSD). (C) Characterize split-initiator HCR suppression.

#### S1.4.3 Measurement of background components

Calculation of the signal-to-background ratio (Section S1.4.2) requires only a Type 1 experiment (using the terminology of Table S5A), yielding the values  $\bar{X}^{\text{SIG+BACK}}$  and  $\bar{X}^{\text{BACK}}$  that are needed to calculate SIG/BACK. If desired, additional control experiments that omit certain reagents can be used to characterize the individual components of background (AF, NSA, NSD). A Type 2 experiment (no probes, hairpins only) yields  $\bar{X}^{\text{NSA+AF}}$  and a Type 3 experiment (no probes, no hairpins) yields  $\bar{X}^{\text{AF}}$ .\* The background components can then be estimated via calculations analogous to (S4) and (S5). The estimated means are:

$$\bar{X}^{\text{NSD}} = \bar{X}^{\text{BACK}} - \bar{X}^{\text{NSA} + \text{AF}} \tag{S8}$$

$$\bar{X}^{\text{NSA}} = \bar{X}^{\text{NSA}+\text{AF}} - \bar{X}^{\text{AF}} \tag{S9}$$

with estimated standard errors:

$$s_{\bar{X}^{\text{NSD}}} \le \sqrt{(s_{\bar{X}^{\text{BACK}}})^2 + (s_{\bar{X}^{\text{NSA}+\text{AF}}})^2}$$
 (S10)

$$s_{\bar{X}NSA} \le \sqrt{(s_{\bar{X}NSA+AF})^2 + (s_{\bar{X}AF})^2}.$$
 (S11)

These upper bounds on estimated standard errors hold under the assumption that the correlations are non-negative for the different components of background. For a given quantity, if the estimated mean is less than the estimated standard error, we report the standard error as an upper bound, and use this bound for uncertainty propagation.

Table S14B provides estimates for AF, NSA, and NSD when imaging *EphA4* in whole-mount chicken embryos using in situ HCR v3.0, all of which are small compared to SIG. Kinetically trapped HCR hairpins automatically suppress NSA and the combination of split-initiator probes and HCR hairpins automatically suppress NSD. Furthermore, HCR generates amplified SIG. All of these factors contribute to achieving a high signal-to-background ratio. If a Type 1 experiment demonstrates SIG  $\gg$  BACK, as is typically the case using in situ HCR v3.0, then there is little motivation to perform Type 2 and Type 3 experiments to characterize the individual background components (AF, NSA, NSD) as these are all bounded above by BACK.

#### S1.4.4 Measurement of split-initiator HCR suppression

To characterize performance of in situ HCR v3.0 in suppressing triggering of HCR by individual split-initiator probes, we augment experiments of Type 1 with additional control experiments that omit certain reagents. First, let us define the NSD and SIG observed using odd probes only or even probes only:

<sup>\*</sup>If a microscope generates non-negligible fluorescence intensities in the absence of sample, this so-called instrument noise (NOISE) should be taken into consideration when calculating background and signal contributions, leading to four Experiment Types (1. SIG+BACK+NOISE, 1. BACK+NOISE, 2. NSA+AF+NOISE, 3. AF+NOISE, 4. NOISE; cf. Table S5AB).

- odd non-specific detection (NSD<sup>odd</sup>): odd probes that bind non-specifically in the sample and subsequently trigger HCR amplification.
- even non-specific detection (NSD<sup>even</sup>): even probes that bind non-specifically in the sample and subsequently trigger HCR amplification.
- odd signal (SIG<sup>odd</sup>): odd probes that hybridize specifically to the target mRNA and subsequently trigger HCR amplification.
- even signal (SIG<sup>even</sup>): even probes that hybridize specifically to the target mRNA in the sample and subsequently trigger HCR amplification.

A Type 4 experiment (odd probes only, with hairpins) yields  $\bar{X}^{BACK^{odd}+SIG^{odd}}$  and  $\bar{X}^{BACK^{odd}}$  and a Type 5 experiment (even probes only, with hairpins) yields  $\bar{X}^{BACK^{even}+SIG^{even}}$  and  $\bar{X}^{BACK^{even}}$ . These quantities in turn can be used to calculate SIG<sup>odd</sup> and SIG<sup>even</sup> via calculations analogous to (S4) and (S5). The estimated means are:

$$\bar{X}^{\text{SIG}^{\text{odd}}} = \bar{X}^{\text{SIG}^{\text{odd}} + \text{BACK}^{\text{odd}}} - \bar{X}^{\text{BACK}^{\text{odd}}}$$
(S12)

$$\bar{X}^{\text{SIGeven}} = \bar{X}^{\text{SIGeven} + \text{BACK}^{\text{even}}} - \bar{X}^{\text{BACK}^{\text{even}}}$$
(S13)

with estimated standard errors:

$$s_{\bar{X}^{\text{SIG}^{\text{odd}}}} \leq \sqrt{(s_{\bar{X}^{\text{SIG}^{\text{odd}}} + \text{BACK}^{\text{odd}})^2 + (s_{\bar{X}^{\text{BACK}^{\text{odd}}}})^2} \tag{S14}$$

$$s_{\bar{X}^{\mathrm{SIG}^{\mathrm{even}}}} \leq \sqrt{(s_{\bar{X}^{\mathrm{SIG}^{\mathrm{even}}}+\mathrm{BACK}^{\mathrm{even}})^2 + (s_{\bar{X}^{\mathrm{BACK}^{\mathrm{even}}}})^2} \tag{S15}$$

These upper bounds on estimated standard errors hold under the assumption that the correlations are non-negative for SIG<sup>odd</sup> and BACK<sup>odd</sup> and for SIG<sup>even</sup> and BACK<sup>even</sup>. For a given quantity, if the estimated mean is less than the estimated standard error, we report the standard error as an upper bound, and use this bound for uncertainty propagation.

Split-initiator HCR suppression can then be characterized by calculating SIG/SIG<sup>odd</sup> and SIG/SIG<sup>even</sup>, with higher values corresponding to more effective suppression. This in situ characterization is akin to the gel studies (Figures 2, S3, and S4) that compare test tube triggering of HCR by odd/even probe pairs colocalized by the target (lane 3) or by either probe alone (lanes 4 or 5). For the in situ data, signal-to-signal ratios are obtained via calculations analogous to (S6) and (S7). The estimated means are:

$$\bar{X}^{\text{SIG/SIG}^{\text{odd}}} = \bar{X}^{\text{SIG}} / \bar{X}^{\text{SIG}^{\text{odd}}}$$
(S16)

$$\bar{X}^{\text{SIG/SIG}^{\text{even}}} = \bar{X}^{\text{SIG}} / \bar{X}^{\text{SIG}^{\text{even}}} \tag{S17}$$

with estimated standard errors:

$$s^{\mathrm{SIG/SIG^{odd}}} \leq \bar{X}^{\mathrm{SIG/SIG^{odd}}} \sqrt{\left(\frac{s_{\bar{X}^{\mathrm{SIG}}}}{\bar{X}^{\mathrm{SIG}}}\right)^2 + \left(\frac{s_{\bar{X}^{\mathrm{SIG^{odd}}}}}{\bar{X}^{\mathrm{SIG^{odd}}}}\right)^2} \tag{S18}$$

$$s^{\mathrm{SIG/SIG^{even}}} \leq \bar{X}^{\mathrm{SIG/SIG^{even}}} \sqrt{\left(\frac{s_{\bar{X}^{\mathrm{SIG}}}}{\bar{X}^{\mathrm{SIG}}}\right)^2 + \left(\frac{s_{\bar{X}^{\mathrm{SIG^{even}}}}}{\bar{X}^{\mathrm{SIG^{even}}}}\right)^2}.$$
(S19)

These upper bounds on estimated standard errors hold under the assumption that the correlations are non-negative for SIG and SIG<sup>odd</sup> and for SIG and SIG<sup>even</sup>.

Table S14C displays the signal-to-signal ratios SIG/SIG<sup>odd</sup> and SIG/SIG<sup>even</sup> when imaging *EphA4* in wholemount chicken embryos using in situ HCR v3.0. Split-initiator probes colocalized by the target are more than an order of magnitude more effective at triggering HCR than odd or even probes alone. Interestingly, with this assay, we are quantifying the automatic *background* suppression capabilities of the split-initiator probes by measuring automatic *signal* suppression, taking advantage of the fact that the target molecules in the embryo will colocalize odd/even probe pairs for a Type 1 experiment and will localize odd or even probes to the same expression region for a Type 4 or 5 experiment.

# S1.4.5 Normalized voxel intensities for qHCR imaging: analog mRNA relative quantitation with subcellular resolution in an anatomical context

For quantitative mRNA imaging using in situ HCR, precision increases with voxel size as long as the imaging voxels remain smaller than the features in the expression pattern (see Section S2.2 of Trivedi *et al.* (2018)). To increase precision, we calculate raw voxel intensities by averaging neighboring pixel intensities while still maintaining a subcellcular voxel size. To facilitate relative quantitation between voxels, we estimate the normalized HCR signal of voxel j in replicate n as:

$$x_{n,j} \equiv \frac{X_{n,j}^{\text{SIG+BACK}} - X^{\text{BOT}}}{X^{\text{TOP}} - X^{\text{BOT}}},$$
(S20)

which translates and rescales the data so that the voxel intensities in each channel fall in the interval [0,1]. Here,

$$X^{\text{BOT}} \equiv \bar{X}^{\text{BACK}} \tag{S21}$$

is the mean background across replicates (see Section S1.4.2) and

$$X^{\text{TOP}} \equiv \max_{n,j} X_{n,j}^{\text{SIG+BACK}}$$
(S22)

is the maximum total fluorescence for a voxel across replicates.

Pairwise expression scatter plots that each display normalized voxel intensities for two channels (e.g., Figures 4 and 5 of Trivedi *et al.* (2018)) provide a powerful quantitative framework for performing multidimensional readout/read-in analyses (Figure 6 of Trivedi *et al.* (2018)). Read-out from anatomical space to expression space enables discovery of expression clusters of voxels with quantitatively related expression levels and ratios (amplitudes and slopes in the expression scatter plots), while read-in from expression space to anatomical space enables discovery of the corresponding anatomical locations of these expression clusters within the embryo. The simple and practical normalization approach of (S20)–(S22) translates and rescales all voxels identically within a given channel (enabling comparison of amplitudes and slopes in scatter plots between replicates), and does not attempt to remove scatter in the normalized signal estimate that is caused by scatter in the background.

To validate relative mRNA quantitation with subcellular resolution ( $2 \times 2 \times 2.7 \mu m$  voxels) in whole-mount chicken embryos, Figures 5C, S18C, and S19C display highly correlated normalized voxel intensities for 2-channel redundant detection of *Dmbx1* and *EphA4*. In this setting, accuracy corresponds to linearity with zero intercept, and precision corresponds to scatter around the line (Trivedi *et al.*, 2018).

# S1.4.6 Dot detection and colocalization for dHCR imaging: digital mRNA absolute quantitation in an anatomical context

To validate the performance of in situ HCR (v3.0) for single-molecule imaging, we perform a 2-channel redundant detection experiment in which a target mRNA is detected using two independent probe sets and HCR amplifiers. Let  $N_1$  denote the number of dots detected in channel 1,  $N_2$  the number of dots detected in channel 2, and  $N_{12}$  the number of colocalized dots appearing in both channels. We define the colocalization fraction for each channel:

$$C_1 = N_{12}/N_1, (S23)$$

$$C_2 = N_{12}/N_2. (S24)$$

As the false-positive and false-negative rates for single-molecule detection go to zero,  $C_1$  and  $C_2$  will both approach 1 from below, providing a quantitative basis for evaluating performance. Colocalization results using in situ HCR v3.0 with split-initiator probes (23-25 probe pairs per channel) in cultured human cells (Table S25) and whole-mount chicken embryos (Table S26) are  $\approx$ 84%, compared with  $\approx$ 50% using in situ HCR v2.0 (39 standard probe pairs per channel) in a previous study in whole-mount zebrafish embryos (Table S27)(Shah *et al.*, 2016).

Single molecules were identified in each channel using the following dot detection algorithm applied to a threedimensional confocal image stack:

- Step 1: Blur noise. To remove noise smaller than the dots of interest, the image was convolved with an isotropic Gaussian blur (standard deviation  $\sigma_{blur}$ ).
- Step 2: Local background subtraction. To eliminate variations in pixel intensity arising from background variations that occur on a length scale larger than the dots of interest, local background subtraction was performed by subtracting the mean pixel intensity of a cube (edge length  $d_{\text{back}}$ ) from the intensity of the pixel at the center of the cube.
- Step 3: Global threshold on pixel intensity. To eliminate dim features, the resulting pixel intensities were subjected to a global threshold ( $t_{\text{pixel}}$ ), and the range [ $t_{\text{pixel}}$ ,1] was renormalized to a [0,1] scale.
- Step 4: Watershed dot detection. To identify single mRNA molecules as dots within the image, regional image maxima were segmented using the minimum saliency watershed method (Couprie & Bertrand, 1997; Yoo *et al.*, 2002). In this method, two maxima are labeled as the same dot if the minimum boundary height between the maxima is less than a given threshold ( $t_{watershed}$ ). Dot coordinates were estimated as the intensity-weighted centroid of each watershed basin, and dot intensities were estimated as the integrated pixel intensity within each basin.
- Step 5: Global threshold on dot intensity. To eliminate dim dots, the resulting dot intensities were normalized on a [0,1] scale and a global threshold  $(t_{dot})$  was applied. The resulting number of dots  $N_i$  was recorded for channel *i*.

After identifying the dots in each channel of a 2-channel redundant detection image, a dot i in Channel 1 and a dot j in Channel 2 were considered colocalized if all of the following four statements were true:

- Test 1: The xy centroids differed by less than a lateral distance threshold ( $r_{xy} = 0.22 \ \mu m$ ).
- Test 2: Dot *i* is the closest dot in Channel 1 to dot *j* in Channel 2.
- Test 3: Dot j is the closest dot in Channel 2 to dot i in Channel 1.
- Test 4: The z centroids differed by less than the axial distance threshold ( $r_z = 0.42 \ \mu m$ ).

Note that due to the lower axial resolution, dot colocalization was tested separately for xy and z. The same distance thresholds were used across all sample types and replicates.<sup>†</sup>

<sup>&</sup>lt;sup>†</sup>For chicken embryo replicate 3, Channels 1 and 2 were manually aligned by applying a constant offset of  $+0.35\mu$ m to the z-coordinates of Channel 1 after the detected dots showed a clear bias in z coordinates between the two channels. No other images were manually aligned.

		HEK cells		Chicken embryos		Zebrafish embryos	
lateral resolution $d_{xy}$ ( $\mu$ m)		0.0624		0.099		0.2167	
axial resolution $d_z$ ( $\mu$ m)		0.42		0.42		0.3369	
Step 1: Gaussian blur radius $\sigma_{\text{blur}}$ ( $\mu$ m)		0.2		0.2		0.1	
Step 2: Mean subtraction cube length $d_{\text{back}} (\mu m)$		2.5		3.0		5.5	
	Replicate	Ch1	Ch2	Ch1	Ch2	Ch1	Ch2
Step 3: Global pixel threshold $(t_{pixel})$	1	0.003	0.012	0.04	0.02	0.02	0.02
	2	0.005	0.012	0.04	0.02	0.01	0.003
	3	0.004	0.01	0.04	0.02	0.01	0.005
Step 4: Watershed saliency minimum ( $t_{watershed}$ )	1	0.15	0.15	0.15	0.15	0.15	0.15
	2	0.15	0.15	0.15	0.15	0.30	0.20
	3	0.15	0.15	0.15	0.15	0.35	0.30
Step 5: Global dot intensity threshold $(t_{dot})$	1	0.01	0.005	0.00	0.01	0.01	0.01
	2	0.02	0.02	0.006	0.013	0.025	0.025
	3	0.03	0.012	0.01	0.025	0.04	0.04

Table S6. Parameters used for dot detection in dHCR images.

# S1.5 Flow cytometry data analysis

Data analysis for flow cytometry experiments on cultured cells closely follows the image analysis of Section S1.4 as detailed below.

## S1.5.1 Raw cell intensities

The components of background (AF, NSA, NSD) and signal (SIG) are defined as before (Section S1.4.1), where n is treated as an index over cells, and i = 1 for each cell since the flow cytometer returns one value per cell.

# S1.5.2 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression for transgenic targets

For a transgenic target mRNA, signal and background are characterized based on flow cytometry experiments of Types 1a and 1b (Table S7A) with SIG+BACK measured in transgenic cells containing the target and BACK measured in wildtype (WT) cells lacking the target. This approach parallels that for characterizing signal and background in images (Section S1.4.2) with transgenic cells taking the place of a region of high expression and WT cells taking the place of a region of no/low expression. If desired, additional control experiments of Types 2 and 3 (Table S7B) can be performed to characterize the components of background (AF, NSA, NSD) using the calculations of Section S1.4.3. Likewise, additional control experiments of Types 4a, 4b, 5a, and 5b (Table S7C) can be performed to characterize split-initiator HCR suppression using the calculations of Section S1.4.4. For transgenic target mR-NAs in human and bacterial cells, Figures S20 and S22 display distributions of cell intensities characterizing signal and background (panel A) and split-initiator HCR suppression (panel B). Tables S17 and S19 display corresponding values for SIG, BACK, and SIG/BACK (panel A), background components AF, NSA, and NSD (panel B), and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>even</sup> (panel C).

			Reage		
	Experiment type	Quantity	Probes	Hairpins	Cell type
Α	1a 1b	SIG+NSD+NSA+AF = SIG+BACK NSD+NSA+AF = BACK	odd + even odd + even	√ √	transgenic WT
В	2 3	NSA+AF AF		$\checkmark$	transgenic transgenic
С	4a 4b 5a 5b	$\begin{split} SIG^{odd} + NSD^{odd} + NSA + AF &= SIG^{odd} + BACK^{odd} \\ NSD^{odd} + NSA + AF &= BACK^{odd} \\ SIG^{even} + NSD^{even} + NSA + AF &= SIG^{even} + BACK^{even} \\ NSD^{even} + NSA + AF &= BACK^{even} \end{split}$	odd odd even even	$\checkmark$	transgenic WT transgenic WT

**Table S7. Experiment types for flow cytometry using in situ HCR v3.0 with a transgenic target mRNA.** (A) Characterize signal, background, and signal-to-background. (B) Characterize components of background (AF, NSA, NSD). (C) Characterize split-initiator HCR suppression.

# S1.5.3 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression for endogenous targets

For an endogenous target mRNA, signal and background are characterized based on flow cytometry experiments of Types 1a and 1b (Table S8A) with SIG+BACK measured using a probe set (odd + even) that address the target in WT cells and BACK measured using a probe set (Tg(odd) and Tg(even)) that addresses a different transgenic target absent from WT cells. Use of a previously validated transgenic probe set to measure background in WT cells ensures that a low measured fluorescence value does not simply indicate a dysfunctional probe set, but indeed represents low background generated by a probe set that is known to be functional if the target is present in the sample. If desired,

Development 145: doi:10.1242/dev.165753: Supplementary information

			Reagents		
	Experiment type	Quantity	Probes	Hairpins	Cell type
Α	1a	SIG+NSD+NSA+AF = SIG+BACK	odd + even	$\checkmark$	WT
	1b	NSD+NSA+AF = BACK	Tg(odd) + Tg(even)	$\checkmark$	WT
В	2	NSA+AF		$\checkmark$	WT
	3	AF			WT
С	4a	$SIG^{odd}$ + $NSD^{odd}$ + $NSA$ + $AF$ = $SIG^{odd}$ + $BACK^{odd}$	odd	$\checkmark$	WT
	4b	$NSD^{odd}$ + $NSA$ + $AF$ = $BACK^{odd}$	Tg(odd)	$\checkmark$	WT
	5a	$SIG^{even} + NSD^{even} + NSA + AF = SIG^{even} + BACK^{even}$	even	$\checkmark$	WT
_	5b	$NSD^{even} + NSA + AF = BACK^{even}$	Tg(even)	$\checkmark$	WT

**Table S8. Experiment types for flow cytometry using in situ HCR v3.0 with an endogenous target mRNA.** (A) Characterize signal, background, and signal-to-background. (B) Characterize components of background (AF, NSA, NSD). (C) Characterize split-initiator HCR suppression. Here, Tg(odd) and Tg(even) denote odd and even probes from a probe set targeting a transgenic mRNA that is absent from WT cells.

additional control experiments of Types 2 and 3 (Table S8B) can be performed to characterize the components of background (AF, NSA, NSD) using the calculations of Section S1.4.3. Likewise, additional control experiments of Types 4a, 4b, 5a, and 5b (Table S8C) can be performed to characterize split-initiator HCR suppression using the calculations of Section S1.4.4. For endogenous target mRNA *GAPDH*, Figure S21 displays distributions of cell intensities characterizing signal and background (panel A) and split-initiator HCR suppression (panel B). Table S18 displays corresponding values for SIG, BACK, and SIG/BACK (panel A), background components AF, NSA, and NSD (panel B), and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>even</sup> (panel C).

# S1.5.4 Normalized single-cell intensities for qHCR flow cytometry: analog mRNA relative quantitation for high-throughput expression profiling of human and bacterial cells

For relative mRNA quantitation between cells, the single-cell intensities within a channel are normalized using equation (S20) with BOT (mean BACK intensity across cells) and TOP (maximum SIG+BACK intensity for a single cell) defined by equations (S21) and (S22). Redundant detection experiments validating mRNA single-cell relative quantitation are displayed for endogenous targets *GAPDH* (Figure S24 and Table S20) and *PGK1* (Figure S25 and Table S21) in HEK cells, and for endogenous target *fusA* in *E. coli* (Figure S26 and Table S22).

# S2 Protocols for in situ HCR v3.0

# S2.1 Protocols for whole-mount chicken embryos

# S2.1.1 Preparation of fixed whole-mount chicken embryos

- 1. Collect chicken embryos on 3M filter paper and place in a petri dish containing Ringer's solution.
- 2. Transfer embryos into a new petri dish with fresh Ringer's solution. NOTE: *This is to rinse away egg yolk before fixation.*
- 3. Transfer into a petri dish containing 4% paraformaldehyde (PFA). CAUTION: Use PFA with extreme care as it is a hazardous material. NOTE: Use fresh PFA to avoid increased autofluorescence.
- 4. Fix the samples at room temperature for 1 h.
- 5. Transfer embryos into a petri dish containing PBST.
- 6. Dissect the embryos off the filter paper. Remove the fixed vitelline membrane (opaque) and cut the squares around area pellucida, without leaving excess of extra-embryonic tissue.
- 7. Transfer embryos into a 2 mL Eppendorf tube containing PBST.
- 8. Nutate for 5 mins with the tube positioned horizontally in a small ice bucket.
- 9. Wash embryos two additional times with 2 mL of PBST, each time nutating for 5 min on ice.
- 10. Dehydrate embryos with  $2 \times 5$  min washes of 2 mL methanol (MeOH) on ice.
- 11. Store embryos at -20 °C overnight before use. NOTE: *Embryos can be stored for six months at -20* °C.
- 12. Transfer the required number of embryos for an experiment to a 2 mL Eppendorf tube. NOTE: *Do not place more than 4 embryos in each 2 mL Eppendorf tube.*
- 13. Rehydrate with a series of graded 2 mL MeOH/PBST washes for 5 min each on ice:
  - (a) 75% MeOH / 25% PBST
  - (b) 50% MeOH / 50% PBST
  - (c) 25% MeOH / 75% PBST
  - (d) 100% PBST
  - (e) 100% PBST.
- 14. Treat embryos with 2 mL of 10 μg/mL proteinase K solution for 2 min (stage HH 8) or 2.5 min (stages HH 10–11) at room temperature.
  NOTE: Proteinase K concentration and treatment time should be reoptimized for each batch of proteinase K, or for samples at a different developmental stage.
- 15. Postfix with 2 mL of 4% PFA for 20 min at room temperature.
- 16. Wash embryos 2  $\times$  5 min with 2 mL of PBST on ice.
- 17. Wash embryos with 2 mL of 50% PBST / 50%  $5\times$  SSCT for 5 min on ice.
- 18. Wash embryos with 2 mL of  $5 \times$  SSCT for 5 min on ice.

# S2.1.2 Buffer recipes for sample preparation

Ringer's solution	For 2 L of solution
123 mM NaCl	14.4 g of NaCl
$1.53 \text{ mM CaCl}_2$	$340 \text{ mg of } CaCl_2$
4.96 mM KCl <sub>2</sub>	740 mg of KCl
$0.81 \text{ mM Na}_2\text{HPO}_4$	$230 \text{ mg of } \text{Na}_2\text{HPO}_4$
0.15 mM KH <sub>2</sub> PO <sub>4</sub>	$40 \text{ mg of } \text{KH}_2 \text{PO}_4$
	Bring volume up to 1.5 L with ultrapure $H_2O$
	Adjust pH to 7.4 and fill up to 2 L with ultrapure $H_2O$
	Filter sterilize with 0.22 $\mu$ m bottle top filter
4% Paraformaldehyde (PFA)	For 25 mL of solution
4% PFA	1 g of PFA powder
$1 \times PBS$	$25 \text{ mL of } 1 \times \text{PBS}$
	Heat solution at 50–60 °C to dissolve powder
PBST	For 50 mL of solution
$\overline{1 \times PBS}$	$\overline{5 \text{ mL of } 10 \times \text{PBS}}$
0.1% Tween 20	500 µL of 10% Tween 20
	Fill up to 50 mL with ultrapure $H_2O$
Proteinase K solution	For 2 mL of solution
$10 \mu\text{g/mL}$ proteinase K	$\frac{1}{\mu L}$ of 20 mg/mL proteinase K
	Fill up to 2 mL with PBST

NOTE: Avoid using calcium chloride and magnesium chloride in PBS as this leads to increased autofluorescence in the embryos.

## S2.1.3 Multiplexed in situ HCR v3.0 using split-initiator probes

#### **Detection stage**

- 1. For each sample, transfer 1-4 embryos to a 2 mL Eppendorf tube. NOTE: *Do not place more than 4 embryos in each 2 mL Eppendorf tube.*
- 2. Incubate embryos in 1 mL of 30% probe hybridization buffer on ice for 5 min. CAUTION: probe hybridization buffer contains formamide, a hazardous material. NOTE: Pre-heat probe hybridization buffer to 37 °C before use.
- 3. Remove the buffer and pre-hybridize with 1 mL of 30% probe hybridization buffer for 30 min at 37 °C.
- Prepare probe solution by adding 4 pmol of each probe mixture (odd & even: 2 μL of 2 μM stock per probe mixture) to 1 mL of 30% probe hybridization buffer at 37 °C.
   NOTE: For Figures 7B and S30 (dHCR), 10 pmol of each probe was used to improve probe hybridization efficiency.
- 5. Remove the pre-hybridization solution and add the probe solution.
- 6. Incubate embryos overnight (12-16 h) at 37 °C.
- 7. Remove excess probes by washing embryos 4 × 15 min with 1 mL of 30% probe wash buffer at 37 °C:
   CAUTION: probe wash buffer contains formamide, a hazardous material.
   NOTE: Pre-heat probe wash buffer to 37 °C before use.
- 8. Wash samples  $2 \times 5$  min with  $5 \times$  SSCT at room temperature.

#### **Amplification stage**

- 1. Pre-amplify embryos with 500 mL of amplification buffer for 5 min at room temperature. NOTE: *Equilibrate amplification buffer to room temperature before use*.
- 2. Prepare 30 pmol of each fluorescently labeled hairpin by snap cooling 10  $\mu$ L of 3  $\mu$ M stock in hairpin storage buffer (heat at 95 °C for 90 seconds and cool to room temperature in a dark drawer for 30 min).
- 3. Prepare hairpin solution by adding all snap-cooled hairpins to 500  $\mu$ L of amplification buffer at room temperature.
- 4. Remove the pre-amplification solution and add the hairpin solution.
- 5. Incubate the embryos overnight (12–16 h) in the dark at room temperature. NOTE: For Figures 7B and S30 (dHCR), a 90 min amplification time was used to ensure single-molecule dots are diffraction-limited.
- 6. Remove excess hairpins by washing with 1 mL of  $5 \times$  SSCT at room temperature:
  - (a)  $2 \times 5 \min$
  - (b)  $2 \times 30 \text{ min}$
  - (c)  $1 \times 5 \min$

# S2.1.4 Buffer recipes for in situ HCR v3.0

Probes, amplifiers, probe hybridization buffer, and probe wash buffer should be stored at -20  $^{\circ}$ C. Amplification buffer should be stored at 4  $^{\circ}$ C. Keep these reagents on ice at all times during probe and amplifier preparation. Make sure all solutions are well mixed before use.

## 30% probe hybridization buffer

30% formamide $5\times$  sodium chloride sodium citrate (SSC)9 mM citric acid (pH 6.0)0.1% Tween 2050 µg/mL heparin $1\times$  Denhardt's solution10% dextran sulfate

## 30% probe wash buffer

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 μg/mL heparin

#### **Amplification buffer**

5× sodium chloride sodium citrate (SSC)0.1% Tween 2010% dextran sulfate

## $5 \times SSCT$

 $5 \times$  sodium chloride sodium citrate (SSC) 0.1% Tween 20

**50% dextran sulfate** 50% dextran sulfate For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin 800  $\mu$ L of 50× Denhardt's solution 8 mL of 50% dextran sulfate Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 20 g of dextran sulfate powder Fill up to 40 mL with ultrapure H<sub>2</sub>O

#### S2.1.5 Sample mounting for microscopy

- 1. Make a chamber for mounting each embryo by aligning two stacks of double-sided tape (2 pieces per stack) 1 cm apart on a 25 mm  $\times$  75 mm glass slide.
- 2. Place an embryo between the tape stacks on the slide and remove as much solution as possible.
- 3. Align the embryo for dorsal imaging and carefully touch the slide with a kimwipe to further dry the area around the embryo.
- 4. Add two drops of SlowFade Gold antifade mountant on top of the embryo.
- 5. Place a 22 mm × 30 mm No. 1 coverslip on top of the stacks to close the chamber. NOTE: See Section S1.2 for details of confocal microscopes used to image whole-mount chicken embryos.

# S2.1.6 Reagents and supplies

Paraformaldehyde (PFA) (Sigma Cat. # P6148) Methanol (Mallinckrodt Chemicals Cat. # 3016-16) Proteinase K, molecular biology grade (NEB Cat. # P8107S) Formamide (Deionized) (Ambion Cat. # AM9342)  $20 \times$  sodium chloride sodium citrate (SSC) (Life Technologies Cat. # 15557-044) Heparin (Sigma Cat. # H3393) 50% Tween 20 (Life Technologies Cat. # 00-3005)  $50 \times$  Denhardt's solution (Life Technologies Cat. # 750018) Dextran sulfate, mol. wt. > 500,000 (Sigma Cat. # D6001)  $25 \text{ mm} \times 75 \text{ mm}$  glass slide (VWR Cat. # 48300-025)  $22 \text{ mm} \times 30 \text{ mm}$  No. 1 coverslip (VWR Cat. # 48393-026) SlowFade Gold antifade mountant (Life Technologies Cat. # S36937)

# S2.2 Protocols for mammalian cells on a chambered slide

# S2.2.1 Preparation of fixed mammalian cells on a chambered slide

- 1. Coat bottom of each chamber by applying 300  $\mu$ L of 0.01% poly-D-lysine prepared in cell culture grade H<sub>2</sub>O. NOTE: A volume of 300  $\mu$ L is sufficient per chamber on an 8-chamber slide.
- 2. Incubate for at least 30 min at room temperature.
- 3. Aspirate the coating solution and wash each chamber twice with molecular biology grade  $H_2O$ .
- 4. Plate desired number of cells in each chamber.
- 5. Grow cells to desired confluency for 24–48 h.
- 6. Aspirate growth media and wash each chamber with 300 μL of DPBS. NOTE: Avoid using calcium chloride and magnesium chloride in DPBS as this leads to increased autofluorescence.
- 7. Add 300 μL of 4% formaldehyde to each chamber.
   CAUTION: Use formaldehyde with extreme care as it is a hazardous material.
- 8. Incubate for 10 min at room temperature.
- 9. Remove fixative and wash each chamber 2  $\times$  300  $\mu L$  of DPBS.
- 10. Aspirate DPBS and add 300  $\mu L$  of ice-cold 70% ethanol.
- 11. Permeabilize cells overnight at -20  $^\circ \text{C}.$
- 12. Cells can be stored at -20  $^\circ C$  or 4  $^\circ C$  until use.

#### S2.2.2 Buffer recipes for sample preparation

4% formaldehyde in PBS

 $\frac{4\% \text{ formaldehyde}}{1 \times \text{PBS}}$ 

 $\label{eq:solution} \begin{array}{l} \hline For \ 10 \ mL \ of \ solution \\ \hline 2.5 \ mL \ of \ 16\% \ formal dehyde \\ 1 \ mL \ of \ 10\times \ PBS \\ \hline Fill \ up \ to \ 10 \ mL \ with \ molecular \ biology \ grade \ H_2O \end{array}$ 

NOTE: Avoid using calcium chloride and magnesium chloride in PBS as this leads to increased autofluorescence in the samples.

## S2.2.3 Multiplexed in situ HCR v3.0 using split-initiator probes

#### **Detection stage**

- 1. Aspirate EtOH and air dry samples at room temperature. NOTE: *Drying of sample is optional.*
- 2. Wash samples two times with 300  $\mu$ L of 2× SSC.
- Pre-hybridize samples in 300 μL of 30% probe hybridization buffer for 30 min at 37 °C. CAUTION: probe hybridization buffer contains formamide, a hazardous material. NOTE: Pre-heat probe hybridization buffer to 37 °C before use.
- Prepare probe solution by adding 1.2 pmol of each probe mixture (odd & even: 0.6 μL of 2 μM stock per probe mixture) to 300 μL of 30% probe hybridization buffer at 37 °C.
   NOTE: For Figures 7A and S29 (dHCR), 3 pmol of each probe was used to improve probe hybridization efficiency.
- 5. Remove the pre-hybridization solution and add the probe solution.
- 6. Incubate samples overnight (12-16 h) at 37 °C.
- 7. Remove excess probes by washing  $4 \times 5$  min with 300  $\mu$ L of 30% probe wash buffer at 37 °C. CAUTION: probe wash buffer contains formamide, a hazardous material. NOTE: Pre-heat probe wash buffer to 37 °C before use.
- 8. Wash samples  $2 \times 5$  min with  $5 \times$  SSCT at room temperature.

#### **Amplification stage**

- 1. Pre-amplify samples in 300  $\mu$ L of amplification buffer for 30 min at room temperature. NOTE: *Equilibrate amplification buffer to room temperature before use*.
- 2. Prepare 18 pmol of each fluorescently labeled hairpin by snap cooling 6  $\mu$ L of 3  $\mu$ M stock in hairpin storage buffer (heat at 95 °C for 90 seconds and cool to room temperature in a dark drawer for 30 min).
- 3. Prepare hairpin solution by adding all snap-cooled hairpins to 300  $\mu$ L of amplification buffer at room temperature.
- 4. Remove the pre-amplification solution and add the hairpin solution.
- 5. Incubate samples overnight (12–16 h) in the dark at room temperature. NOTE: For Figures 7A and S29 (dHCR), a 45 min amplification time was used to ensure single-molecule dots are diffraction-limited.
- 6. Remove excess hairpins by washing 5  $\times$  5 min with 300  $\mu$ L of 5  $\times$  SSCT at room temperature.
- 7. Aspirate  $5 \times$  SSCT and add  $\approx 100 \ \mu$ L of SlowFade Gold antifade mountant with DAPI.
- 8. Samples can be stored at 4 °C protected from light prior to imaging.

# S2.2.4 Buffer recipes for in situ HCR v3.0

Probes, amplifiers, probe hybridization buffer, and probe wash buffer should be stored at -20  $^{\circ}$ C. Amplification buffer should be stored at 4  $^{\circ}$ C. Keep these reagents on ice at all times during probe and amplifier preparation. Make sure all solutions are well mixed before use.

## 30% probe hybridization buffer

30% formamide $5\times$  sodium chloride sodium citrate (SSC)9 mM citric acid (pH 6.0)0.1% Tween 2050  $\mu$ g/mL heparin1× Denhardt's solution10% dextran sulfate

## 30% probe wash buffer

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 μg/mL heparin

#### **Amplification buffer**

5× sodium chloride sodium citrate (SSC)0.1% Tween 2010% dextran sulfate

## $5 \times SSCT$

 $5 \times$  sodium chloride sodium citrate (SSC) 0.1% Tween 20

**50% dextran sulfate** 50% dextran sulfate For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin 800  $\mu$ L of 50× Denhardt's solution 8 mL of 50% dextran sulfate Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 20 g of dextran sulfate powder Fill up to 40 mL with ultrapure H<sub>2</sub>O

# S2.2.5 Reagents and supplies

Molecular biology grade H<sub>2</sub>O (Corning Cat. # 46-000-CV) 16% Formaldehyde (w/v), Methanol-free (Life Technologies Cat. # 28906) DPBS, no calcium, no magnesium (Life Technologies Cat. # 14190144) 10× PBS (Ambion Cat. # AM9624) Formamide (Deionized) (Ambion Cat. # AM9342) 20× sodium chloride sodium citrate (SSC) (Life Technologies Cat. # 15557-044) Heparin (Sigma Cat. # H3393) 10% Tween 20 (BioRad Cat. # 161-0781) 50× Denhardt's solution (Life Technologies Cat. # 750018) Dextran sulfate, mol. wt. > 500,000 (Sigma Cat. # D6001) ibidi  $\mu$ -slide ibitreat (ibidi Cat. # 80826) SlowFade Gold antifade mountant with DAPI (Life Technologies Cat. # S36938)

## S2.3 Protocols for mammalian cells in suspension

#### S2.3.1 Preparation of fixed mammalian cells in suspension

- 1. Aspirate growth media from culture plate and wash cells with DPBS. NOTE: Avoid using calcium chloride and magnesium chloride in DPBS as this leads to increased autofluorescence.
- 2. Add 3 mL of trypsin per 10 cm plate and incubate in a 5% CO<sub>2</sub> incubator for 5 min at 37 °C.
- 3. Quench trypsin by adding 3 mL of growth media.
- 4. Transfer cells to a conical tube and centrifuge for 5 min at  $180 \times g$ .
- 5. Aspirate supernatant and re-suspend cells in 4% formaldehyde to reach  $\approx 10^6$  cells/mL. CAUTION: Use formaldehyde with extreme care as it is a hazardous material.
- 6. Fix cells for 1 hr at room temperature.
- 7. Centrifuge for 5 min at  $180 \times g$  and remove supernatant.
- 8. Wash cells 4 times with PBST (use the same volume as formaldehyde solution). NOTE: Centrifuge for 5 min at  $180 \times g$  and aspirate supernatant between washes.
- 9. Re-suspend cells in ice-cold 70% ethanol (use the same volume as formaldehyde and PBST solutions).
- 10. Permeabilize cells overnight at 4  $^{\circ}$ C.
- 11. Cells can be stored at 4 °C until use.

#### S2.3.2 Buffer recipes for sample preparation

**4% formaldehyde in PBST** 4% formaldehyde

 $1 \times PBS$ , 0.1% Tween 20

For 36 mL of solution 9 mL of 16% formaldehyde 3.6 mL of 10× PBST 180  $\mu$ L of 10% Tween 20 Fill up to 36 mL with ultrapure H<sub>2</sub>O

NOTE: Avoid using calcium chloride and magnesium chloride in PBS as this leads to increased autofluorescence in the samples.

## S2.3.3 Multiplexed in situ HCR v3.0 using split-initiator probes

#### **Detection stage**

- 1. Transfer desired amount  $(0.5-1 \times 10^6)$  of fixed cells into a 1.5 mL Eppendorf tube.
- Centrifuge for 5 min to remove EtOH. NOTE: All centrifugation steps are done at 180 × g.
- 3. Wash cells twice with 500  $\mu$ L of PBST. Centrifuge for 5 min to remove supernatant.
- 4. Re-suspend the pellet with 400 μL of 30% LMW probe hybridization buffer and pre-hybridize for 30 min at 37 °C.
  CAUTION: probe hybridization buffer contains formamide, a hazardous material.
  NOTE: Pre-heat probe hybridization buffer to 37 °C before use.
- 5. In the meantime, prepare probe solution by adding 2 pmol of each probe mixture (odd & even: 1  $\mu$ L of 2  $\mu$ M stock per probe mixture) to 100  $\mu$ L of 30% LMW probe hybridization buffer at 37 °C.
- 6. Add the probe solution directly to the sample to reach a final probe concentration of 4 nM.
- 7. Incubate the sample overnight at 37  $^{\circ}$ C.
- 8. Centrifuge for 5 min to remove probe solution.
- 9. Re-suspend the cell pellet with 500 μL of 30% probe wash buffer.
   CAUTION: probe wash buffer contains formamide, a hazardous material.
   NOTE: Pre-heat probe wash buffer to 37 °C before use.
- 10. Incubate for 10 min at 37 °C and remove the wash solution by centrifugation for 5 min.
- 11. Repeat steps 9 and 10 for three additional times.
- 12. Re-suspend the cell pellet with 500  $\mu$ L of 5× SSCT.
- 13. Incubate for 5 min at room temperature.
- 14. Proceed to hairpin amplification.

#### **Amplification stage**

- 1. Centrifuge for 5 min to pellet the cells.
- Re-suspend the cell pellet with 150 μL of LMW amplification buffer and pre-amplify for 30 min at room temperature.
   NOTE: Equilibrate amplification buffer to room temperature before use.
- 3. Prepare 15 pmol of each fluorescently labeled hairpin by snap cooling 5  $\mu$ L of 3  $\mu$ M stock in hairpin storage buffer (heat at 95 °C for 90 seconds and cool to room temperature in a dark drawer for 30 min).
- 4. Prepare hairpin mixture by adding all snap-cooled hairpins to 100  $\mu$ L of LMW amplification buffer at room temperature.
- 5. Add the hairpin mixture directly to the sample to reach a final hairpin concentration of 60 nM.
- 6. Incubate the sample overnight (>12 h) in the dark at room temperature.
- 7. Centrifuge for 5 min and remove the hairpin solution.

- 8. Re-suspend the cell pellet with 500  $\mu$ L of 5× SSCT.
- 9. Without incubation, remove the wash solution by centrifugation for 5 min.
- 10. Repeat steps 8 and 9 for five additional times.
- 11. Re-suspend the cell pellet in desired buffer and volume. NOTE: Samples can be stored at 4 °C protected from light before flow cytometry or imaging.
- 12. Filter cells before flow cytometry.

# S2.3.4 Buffer recipes for in situ HCR v3.0

Probes, amplifiers, probe hybridization buffer, and probe wash buffer should be stored at -20  $^{\circ}$ C. Amplification buffer should be stored at 4  $^{\circ}$ C. Keep these reagents on ice at all times during probe and amplifier preparation. Make sure all solutions are well mixed before use.

#### 30% probe hybridization buffer (LMW d.s.)

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 μg/mL heparin
1× Denhardt's solution
10% low MW dextran sulfate

## 30% probe wash buffer

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 μg/mL heparin

## Amplification buffer (LMW d.s.)

5× sodium chloride sodium citrate (SSC)0.1% Tween 2010% low MW dextran sulfate

# 5× SSCT

 $5 \times$  sodium chloride sodium citrate (SSC) 0.1% Tween 20

# $\underline{1 \times PBST}$

1× PBS 0.1% Tween 20

# 50% dextran sulfate

50% dextran sulfate

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin 800  $\mu$ L of 50× Denhardt's solution 8 mL of 50% low MW dextran sulfate Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin Fill up to 40 mL with ultrapure H<sub>2</sub>O

 $\label{eq:solution} \begin{array}{l} \hline \mbox{For 40 mL of solution} \\ 10 \mbox{ mL of 20} \times \mbox{SSC} \\ 400 \mbox{ $\mu$L of 10\% Tween 20} \\ 8 \mbox{ mL of 50\% low MW dextran sulfate} \\ \hline \mbox{Fill up to 40 mL with ultrapure } H_2O \end{array}$ 

 $\label{eq:solution} \begin{array}{l} \hline \mbox{For 40 mL of solution} \\ 10 \mbox{ mL of 20} \times \mbox{ SSC} \\ 400 \mbox{ $\mu$L of 10\% Tween 20} \\ \mbox{fill up to 40 mL with ultrapure } \mbox{ H}_2\mbox{O} \end{array}$ 

 $\frac{\text{For 40 mL of solution}}{10 \text{ mL of } 10 \times \text{PBST (0.5\% Tween 20)}}$  $200 \ \mu\text{L of 10\% Tween 20}$ Fill up to 40 mL with ultrapure H<sub>2</sub>O

 $\frac{For \ 40 \ mL \ of \ solution}{20 \ g \ of \ low \ MW \ dextran \ sulfate \ powder}$  Fill up to 40 mL with ultrapure  $H_2O$ 

# S2.3.5 Reagents and supplies

DPBS, no calcium, no magnesium (Life Technologies Cat. # 14190144) Trypsin-EDTA (0.25%), phenol red (Life Technologies Cat. # 25200072) 16% Formaldehyde (w/v), methanol-free (Life Technologies Cat. # 28908) 10× PBST (Rockland Cat. # MB-075-1000) 10% Tween 20 solution (Bio-Rad Cat. # 161-0781) Formamide (Deionized) (Ambion Cat. # AM9342) 20× sodium chloride sodium citrate (SSC) (Life Technologies Cat. # 15557-044) Heparin (Sigma Cat. # H3393) 50% Tween 20 (Life Technologies Cat. # 00-3005) 50× Denhardt's solution (Life Technologies Cat. # 750018) Dextran sulfate, mol. wt. 6,500-10,000 (Sigma Cat. # D4911)

## S2.4 Protocols for bacteria in suspension

#### S2.4.1 Preparation of fixed bacteria in suspension

- 1. Grow *E. coli* from streaked plate or frozen glycerol stocks in 2–3 mL of LB media overnight in a 37 °C shaker.
- 2. Dilute to make a 5 mL liquid culture with  $OD_{600} = 0.05$ .
- 3. Incubate in a 37 °C shaker until  $OD_{600} \approx 0.5$  (exponential phase).
- Aliquot 1 mL of cells and centrifuge for 10 min.
   NOTE: Centrifugation should be a gentle as possible to pellet cells. For E. coli all centrifugation steps are done at 4000 × g.
- 5. Remove supernatant and re-suspend cells in 750  $\mu$ L 1 × phosphate-buffered saline (PBS). NOTE: *Remove all solutions via pipetting throughout the protocol.*
- 6. Add 250  $\mu$ L of 4% formaldehyde to and incubate overnight at 4 °C. CAUTION: Use formaldehyde with extreme care as it is a hazardous material.
- 7. Centrifuge for 10 min and remove supernatant.
- 8. Re-suspend cells in 150  $\mu$ L 1 × PBS.
- 9. Add 850  $\mu$ L of 100% MeOH and store at -20 °C before use.
## S2.4.2 Buffer recipes for sample preparation

## LB media

5 g of Novagen LB Broth Miller powder Fill up to 200 mL with ultrapure  $H_2O$  Autoclave at 121 °C for 20 min

#### **4% formaldehyde in PBS 4% formaldehyde**

 $1 \times PBS$ 

 $\label{eq:entropy} \begin{array}{l} \hline For \ 4 \ mL \ of \ solution \\ 1 \ mL \ of \ 16\% \ formal dehyde \\ 0.4 \ mL \ of \ 10\times \ PBS \\ \hline Fill \ up \ to \ 4 \ mL \ with \ ultrapure \ H_2O \end{array}$ 

NOTE: Avoid using calcium chloride and magnesium chloride in PBS as this leads to increased autofluorescence in the samples.

### S2.4.3 Multiplexed in situ HCR v3.0 using split-initiator probes

#### **Detection stage**

- 1. Transfer 150  $\mu$ L of fixed cells into a 1.5 mL Eppendorf tube.
- 2. Centrifuge for 5 min and remove supernatant.
- 3. Wash cells with 500  $\mu$ L of 1× PBST. Centrifuge for 5 min to remove supernatant.
- 4. Re-suspend the pellet with 400 μL of 30% LMW probe hybridization buffer and pre-hybridize for 1 hr at 37 °C.
  CAUTION: probe hybridization buffer contains formamide, a hazardous material.
  NOTE: Pre-heat probe hybridization buffer to 37 °C before use.
- 5. In the meantime, prepare probe solution by adding 2 pmol of each probe mixture (odd & even: 1  $\mu$ L of 2  $\mu$ M stock per probe mixture) to 100  $\mu$ L of 30% LMW probe hybridization buffer at 37 °C.
- 6. Add the probe solution directly to the sample to reach a final probe concentration of 4 nM.
- 7. Incubate the sample overnight at 37  $^{\circ}$ C.
- 8. Add 1mL of probe wash buffer to the sample.
   CAUTION: probe wash buffer contains formamide, a hazardous material.
   NOTE: Pre-heat probe wash buffer to 37 °C before use.
- 9. Centrifuge for 5 min and remove the wash solution.
- 10. Re-suspend the cell pellet with 500  $\mu$ L wash solution.
- 11. Incubate for 5 min at 37 °C and remove the wash solution by centrifugation for 5 min.
- 12. Repeat steps 10 and 11 for two additional times but with 10 min incubation.
- 13. Proceed to hairpin amplification.

#### **Amplification stage**

1. Re-suspend the cell pellet with 150  $\mu$ L of LMW amplification buffer and pre-amplify for 30 min at room temperature.

NOTE: Equilibrate amplification buffer to room temperature before use.

- 2. Prepare 15 pmol of each fluorescently labeled hairpin by snap cooling 5  $\mu$ L of 3  $\mu$ M stock in hairpin storage buffer (heat at 95 °C for 90 seconds and cool to room temperature in a dark drawer for 30 min).
- 3. Prepare hairpin mixture by adding all snap-cooled hairpins to 100  $\mu$ L of LMW amplification buffer at room temperature.
- 4. Add the hairpin mixture directly to the sample to reach a final hairpin concentration of 60 nM.
- 5. Incubate the sample overnight (>12 h) in the dark at room temperature.
- 6. Add 1 mL of  $5 \times$  SSCT at room temperature to the sample to dilute the solution.
- 7. Centrifuge for 5 min and remove the hairpin solution.
- 8. Re-suspend the cell pellet with 500  $\mu$ L of 5× SSCT and incubate 5 min at room temperature.
- 9. Centrifuge for 5 min and remove the wash solution.

- 10. Repeat steps 8 and 9 for two additional times but with a 10 min incubation.
- 11. Re-suspend the cell pellet in desired buffer and volume. NOTE: Samples can be stored at 4 °C protected from light before flow cytometry.
- 12. Filter cells before flow cytometry.

### S2.4.4 Buffer recipes for in situ HCR v3.0

Probes, amplifiers, probe hybridization buffer, and probe wash buffer should be stored at -20 °C. Amplification buffer should be stored at 4 °C. Make sure all solutions are well mixed before use.

#### 30% probe hybridization buffer (low MW D. S.)

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 µg/mL heparin
1× Denhardt's solution
10% low MW dextran sulfate

#### 30% probe wash buffer

30% formamide
5× sodium chloride sodium citrate (SSC)
9 mM citric acid (pH 6.0)
0.1% Tween 20
50 μg/mL heparin

### Amplification buffer (low MW D. S.)

5× sodium chloride sodium citrate (SSC)0.1% Tween 2010% low MW dextran sulfate

## 5× SSCT

 $5 \times$  sodium chloride sodium citrate (SSC) 0.1% Tween 20

 $\frac{1 \times PBST}{1 \times PBS}$ 0.1% Tween 20

### 50% dextran sulfate

50% dextran sulfate

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin 800  $\mu$ L of 50× Denhardt's solution 8 mL of 50% low MW dextran sulfate Fill up to 40 mL with ultrapure H<sub>2</sub>O

For 40 mL of solution 12 mL formamide 10 mL of  $20 \times$  SSC 360  $\mu$ L 1 M citric acid, pH 6.0 400  $\mu$ L of 10% Tween 20 200  $\mu$ L of 10 mg/mL heparin Fill up to 40 mL with ultrapure H<sub>2</sub>O

 $\label{eq:solution} \begin{array}{l} \hline For \ 40 \ \text{mL of solution} \\ 10 \ \text{mL of } 20 \times \ \text{SSC} \\ 400 \ \mu\text{L of } 10\% \ \text{Tween } 20 \\ 8 \ \text{mL of } 50\% \ \text{low MW dextran sulfate} \\ \hline Fill \ \text{up to } 40 \ \text{mL with ultrapure } H_2O \end{array}$ 

 $\frac{\text{For 40 mL of solution}}{10 \text{ mL of } 10 \times \text{PBST} (0.5\% \text{ Tween 20})}$   $200 \ \mu\text{L of } 10\% \text{ Tween 20}$ Fill up to 40 mL with ultrapure H<sub>2</sub>O

 $\frac{For \ 40 \ mL \ of \ solution}{20 \ g \ of \ low \ MW \ dextran \ sulfate \ powder}$ Fill up to 40 mL with ultrapure H<sub>2</sub>O

## S2.4.5 Reagents and supplies

LB Broth Miller (Novagen Cat. # 71753-5) 16% Formaldehyde (w/v), methanol-free (Life Technologies Cat. # 28908) 10× PBS (Ambion Cat. # AM9624) 10× PBST (Rockland Cat. # MB-075-1000) 10% Tween 20 solution (Bio-Rad Cat. # 161-0781) Formamide (Deionized) (Ambion Cat. # AM9342) 20× sodium chloride sodium citrate (SSC) (Life Technologies Cat. # 15557-044) Heparin (Sigma Cat. # H3393) 50% Tween 20 (Life Technologies Cat. # 00-3005) 50× Denhardt's solution (Life Technologies Cat. # 750018) Dextran sulfate, mol. wt. 6,500-10,000 (Sigma Cat. # D4911)

## S3 Additional studies

## S3.1 Validation of split-initiator HCR suppression in vitro and in situ (cf. Figure 2)

Figures S3 and S4 display gel studies measuring split-initiator HCR suppression for amplifiers B1–B5, revealing typical  $\approx$ 60-fold suppression. Table S9 displays signal-to-signal ratios for the same amplifiers used in situ within whole-mount chicken embryos (imaging) and/or within cultured human or bacterial cells (flow cytometry), revealing typical  $\approx$ 50-fold suppression.



Figure S3. Test tube validation of split-initiator HCR suppression for amplifiers B1, B2, B4, and B5 (cf. Figure 2). (A) Agarose gel electrophoresis. Reaction conditions: hairpins H1 and H2 at 0.5  $\mu$ M each (Lanes 1-7); DNA oligos I1, P1, P2, and/or Target at 5 nM each (lanes noted on the gel); 5× SSCT buffer; overnight reaction at room temperature. Hairpins H1 and H2 labeled with Alexa 647 fluorophore (green channel). dsDNA 1 kb ladder pre-stained with SYBR Gold (red channel). (B) Quantification of the polymer band in panel (A).

Organism	Target	Samples	Channel	$SIG/SIG^{odd}$	SIG/SIG <sup>even</sup>	Table
G. gallus domesticus	EphA4	3 embryos	B2-Alexa647	>800	$57 \pm 5$	S14
H. sapiens sapiens	Tg(d2eGFP)	55,000 cells	B3-Alexa594	$461 \pm 9$	$22.2\pm0.1$	S17
H. sapiens sapiens	GAPDH	30,000 cells	B4-Alexa594	$55 \pm 5$	$42.9\pm0.6$	S18
H. sapiens sapiens	GAPDH	20,000 cells	B5-Alexa488	$40.4\pm0.8$	>3000	S20
H. sapiens sapiens	GAPDH	20,000 cells	B4-Alexa594	$67 \pm 2$	$52 \pm 1$	S20
H. sapiens sapiens	PGK1	54,000 cells	B1-Alexa488	>5000	$49.0\pm0.5$	S21
H. sapiens sapiens	PGK1	54,000 cells	B2-Alexa594	$41 \pm 1$	$13 \pm 1$	S21
H. sapiens sapiens	GAPDH	18,000 cells	B4-Alexa488	$93 \pm 6$	$91 \pm 14$	S23
H. sapiens sapiens	PGK1	18,000 cells	B2-Alexa594	$21 \pm 2$	$18.8\pm0.4$	S23
E. coli	Tg(eGFP)	18,000 cells	B3-Alexa594	>3000	$9.9\pm0.5$	S19
E. coli	fusA	3,400 cells	B3-Alexa488	$40 \pm 20$	$14 \pm 3$	S22
E. coli	fusA	3,400 cells	B2-Alexa594	$50 \pm 20$	$6.2\pm0.6$	S22
E. coli	fusA	35,000 cells	B3-Alexa488	$600 \pm 300$	$17.2\pm1.3$	S24
E. coli	icd	35,000 cells	B1-Alexa594	$800 \pm 300$	$85 \pm 4$	S24

Table S9. In situ validation of split-initiator HCR suppression.



Figure S4. Test tube validation of split-initiator HCR suppression for amplifier B3 (cf. Figure 2). (A) Agarose gel electrophoresis (Replicate 1 is displayed in Figure 2). Reaction conditions: hairpins H1 and H2 at 0.5  $\mu$ M each (Lanes 1-7); DNA oligos I1, P1, P2, and/or Target at 5 nM each (lanes noted on the gel); 5× SSCT buffer; overnight reaction at room temperature. Hairpins H1 and H2 labeled with Alexa 647 fluorophore (green channel). dsDNA 1 kb ladder pre-stained with SYBR Gold (red channel). (B) Quantification of the polymer band in panel (A).

## S3.2 In situ validation of automatic background suppression with split-initiator probes in wholemount chicken embryos (cf. Figure 3)

The following studies are included:

- Measurement of background and signal-to-background for unoptimized standard and split-initiator probe sets as a function of probe set size. Probe sets with 5, 10, or 20 standard probes or split-initiator probe pairs. For standard probes, the probe set size is increased from 5 to 10 to 20 by adding untested probes to a previously validated set of 5 probes Choi *et al.* (2016); each standard probe has a 50-nt target-binding site and carries two HCR initiators (one at each end). Each split-initiator probe pair addresses the nearly identical target subsequence as the corresponding standard probe; each split-initiator probe has a 25-nt target-binding site and carries half an HCR initiator; the two probes within a pair are separated by 2 nt along the target so the overall target binding site for a probe pair is 52 nt. For background comparisons (Figures S5 and S6 and Table S10), the PMT gain is held constant for all experiments to enable comparison of background intensities between experiments. For signal-to-background comparisons (Figures S7 and S8 and Table S11), the PMT gain is adjusted to use the full dynamic range for each probe set.
- Measurement of background and signal-to-background for standard and split-initiator probes with identical target-binding domains. For these studies, standard probes are constructed from split-initiator probe pairs as follows. For each split-initiator probe pair in a probe set, the full initiator is shifted onto either the odd probe (full-initiator odd probe + initiator-free even probe) or onto the even probe (initiator-free odd probe + full-initiator even probe). Within each pair, one probe is then a full-initiator standard probe and one probe is a helper probe that contains no initiator. The helper probes are employed to ensure that each standard probe pair (full-initiator probe + helper probe) has the same target-binding capabilities as its analogous split-initiator probe pair. Probe sets with 20 standard probe pairs or split-initiator probe pairs. For background comparisons (Figure S9 and Table S12), the PMT gain is held constant for all experiments to enable comparison of background intensities between experiments. For signal-to-background comparisons (Figure S10 and Table S13), the PMT gain is adjusted to use the full dynamic range for each probe set.
- Measurement of signal, background, and signal-to-background, background components, and splitinitiator HCR suppression. Measurement of signal, background, and signal-to-background (Figure S11A and Table S14A) using the methods of Section S1.4.2. Measurement of background components (AF, NSA, NSD; Figure S11B and Table S14B) using the methods of Section S1.4.3. Measurement of split-initiator HCR suppression (Figure S11C and Table S14C) using the methods of Section S1.4.4.

## S3.2.1 Measurement of background and signal-to-background for unoptimized standard and split-initiator probe sets as a function of probe set size



**Figure S5. Measurement of background for unoptimized standard probe sets as a function of probe set size (cf. Figure 3A).** Confocal images collected with the microscope PMT gain adjusted to avoid saturating pixels using 20 standard probes (row 3). Protocol: in situ HCR v2.0 (Section S8 of Choi *et al.* (2016)). Target mRNA: *Sox10*. Probe set: 5, 10, or 20 standard probes. Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 11.



**Figure S6.** Measurement of background for unoptimized split-initiator probe sets as a function of probe set size (cf. **Figure 3A).** Confocal images collected with the microscope PMT gain adjusted to avoid saturating pixels using 20 standard probes (Figure S5, row 3). Protocol: in situ HCR v3.0 (Section S2.1). Target mRNA: *Sox10*. Probe set: 5, 10, or 20 split-initiator probe pairs. Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 11.

Probe type	Probe set size	BACK
Standard (v2.0)	5	$210\pm30$
	10	$1160 \pm 60$
	20	$1500\pm100$
Split-initiator (v3.0)	5	$29 \pm 1$
	10	$26 \pm 3$
	20	$28\pm1$

Table S10. Estimated background for standard and split-initiator probes sets as a function of probe set size (cf. Figure 3A). Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figures S5 and S6 using methods of Section S1.4.2.



**Figure S7.** Measurement of signal and background for unoptimized standard probe sets as a function of probe set size (cf. Figure 3B). (A) Confocal images collected with the microscope PMT gain optimized for each probe set. (B) Pixel intensity histograms for Signal + Background (pixels within solid boundary) and Background (pixels within dashed boundary) per embryo in panel A. Protocol: in situ HCR v2.0 (Section S8 of Choi *et al.* (2016)). Target mRNA: *Sox10*. Probe set: 5, 10, or 20 standard probes. Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 11.



**Figure S8.** Measurement of signal and background for unoptimized split-initiator probe sets as a function of probe set size (cf. Figure 3B). (A) Confocal images collected with the microscope PMT gain optimized for each probe set. (B) Pixel intensity histograms for Signal + Background (pixels within solid boundary) and Background (pixels within dashed boundary) per embryo in panel A. Protocol: in situ HCR v3.0 (Section S2.1). Target mRNA: *Sox10*. Probe set: 5, 10, or 20 split-initiator probe pairs. Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 11.

Probe type	Probe set size	BACK	SIG+BACK	SIG	SIG/BACK
Standard (v2.0)	5 10 20	$\begin{array}{c} 200.3 \pm 1.1 \\ 540 \ \pm 40 \\ 720 \ \pm 10 \end{array}$	$1770 \pm 70$ $2170 \pm 40$ $2200 \pm 200$	$1570 \pm 70 \\ 1630 \pm 60 \\ 1400 \pm 200$	$\begin{array}{c} 7.8 \pm 0.3 \\ 3.0 \pm 0.2 \\ 2.0 \pm 0.3 \end{array}$
Split-initiator (v3.0)	5 10 20	$\begin{array}{c} 43.7 \pm 0.6 \\ 34.0 \pm 1.2 \\ 37 \ \pm 3 \end{array}$	$1270 \pm 70$ $1480 \pm 30$ $2460 \pm 40$	$1230 \pm 70$ $1450 \pm 30$ $2420 \pm 40$	$28.2 \pm 1.7 \\ 42.5 \pm 1.8 \\ 65 \pm 5$

Table S11. Estimated signal-to-background for standard and split-initiator probes sets as a function of probe set size (cf. Figure 3B). For this signal-to-background study, the microscope PMT gain was optimized for each probe set so raw background and signal values should only be compared within row. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figures S7 and S8 using methods of Section S1.4.2.

## S3.2.2 Measurement of background and signal-to-background for standard and split-initiator probes with identical target-binding domains



**Figure S9.** Measurement of background for standard and split-initiator probes with identical target-binding domains. (A) Confocal images collected with the microscope PMT gain adjusted to avoid saturating pixels using full-initiator odd probe sets (top row). Protocol: in situ HCR v3.0 (Section S2.1). Target mRNA: *Sox10*. Probe sets (20 probe pairs per probe set): full-initiator odd probes + initiator-free even probes (top row), initiator-free odd probes + full-initiator even probes (middle row), split-initiator odd and even probes (bottom row). Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 10.

Sample	BACK	
Full-initiator probes (odd) Full-initiator probes (even) Split-initiator probes	$\begin{array}{rrr} 910 & \pm  80 \\ 330 & \pm  40 \\ 19.9 \pm 0.9 \end{array}$	

Table S12. Estimated background for standard and split-initiator probes with identical target-binding domains. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figure S9 using methods of Section S1.4.2.



**Figure S10.** Measurement of signal and background for standard and split-initiator probes with identical target-binding domains. (A) Confocal images collected with the microscope PMT gain optimized for each probe set. (B) Pixel intensity histograms for Signal + Background (pixels within solid boundary) and Background (pixels within dashed boundary) per embryo in panel A. Protocol: in situ HCR v3.0 (Section S2.1). Target mRNA: *Sox10*. Probe sets (20 probe pairs per probe set): full-initiator odd probes + initiator-free even probes (top row), initiator-free odd probes + full-initiator even probes (middle row), split-initiator odd and even probes (bottom row). Amplifier: B3-Alexa647. Whole-mount chicken embryos fixed stage HH 10.

Sample	BACK	SIG+BACK	SIG	SIG/BACK
Full-initiator probes (odd)	910 $\pm 80$	$1230 \pm 110$	$320\pm140$	$0.4 \pm 0.2$
Full-initiator probes (even)	$330 \pm 40$	$1610\pm150$	$1280 \pm 160$	$3.9\pm0.6$
Split-initiator probes	$26.4\pm0.7$	$1870 \pm 110$	$1840 \pm 110$	$70 \pm 5$

Table S13. Estimated signal-to-background for standard and split-initiator probes with identical target-binding domains. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figure S10 using methods of Section S1.4.2.

## S3.2.3 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression



**Figure S11. Measurement of signal and background, background components, and split-initiator HCR suppression.** Confocal image: fluorescence merged with bright field. (A) Signal and background: use experiment of Type 1 in Table S5A (odd probes + even probes + hairpins) to measure SIG+BACK (region of high expression) and BACK (region of no/low expression). (B) Background components: use experiment of Type 2 in Table S5B (no probes, hairpins only) to measure NSA+AF (region of high expression); use experiment of Type 3 (no probes, no hairpins) to measure AF (region of high expression). (C) Split-initiator HCR suppression: use experiment of Type 4 in Table S5C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (region of high expression); use experiment of Type 4 in Table S5C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (region of high expression) and BACK<sup>odd</sup> (region of no/low expression); use experiment of Type 5 in Table S5C (even probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (region of high expression) and BACK<sup>even</sup> (region of no/low expression). Left: confocal images collected with the microscope PMT gain optimized to avoid saturating pixels using the full method (top row). Right: pixel intensity histograms for a region of high expression (pixels within solid boundary) and/or low/no expression (pixels within dashed boundary) per embryo. Protocol: in situ HCR v3.0 (Section S2.1). Target mRNA: *EphA4*. Probe set: 20 split-initiator probe pairs. Amplifier: B2-Alexa647. Whole-mount chicken embryos fixed stage HH 11. Development 145: doi:10.1242/dev.165753: Supplementary information

		Ch	annel	Reage	ents	Expression	
	Quantity	B2-A	lexa647	Probes	Hairpins	region	Figure
A	SIG+NSD+NSA+AF = SIG+BACK	1260	$\pm 30$	odd + even	<b>√</b>	high	S11C
	NSD+NSA+AF = BACK	19	$\pm 1$	odd + even	$\checkmark$	low/no	S11C
	SIG	1240	$\pm 30$				
	SIG/BACK	67	$\pm 4$				
В	NSA+AF	8.1	$\pm 0.9$		$\checkmark$	high	S11B
	AF	6.97	$7 \pm 0.08$			high	S11A
	NSA	1.2	$\pm 0.9$				
	NSD	10.5	$\pm 1.3$				
С	$NSD^{odd}$ + $NSA$ + $AF$ = $BACK^{odd}$	11.0	$\pm 1.0$	odd	$\checkmark$	low/no	S11D
	$NSD^{even}$ + $NSA$ + $AF$ = $BACK^{even}$	10.0	$\pm 0.6$	even	$\checkmark$	low/no	S11E
	SIG <sup>odd</sup> +BACK <sup>odd</sup>	11.5	$\pm 1.1$	odd	$\checkmark$	high	S11D
	SIG <sup>even</sup> +BACK <sup>even</sup>	31.8	$\pm 1.7$	even	$\checkmark$	high	S11E
	NSD <sup>odd</sup>	2.8	$\pm 1.4$				
	NSD <sup>even</sup>	1.9	$\pm 1.1$				
	SIG <sup>odd</sup>	$<\!1.5$					
	SIG <sup>even</sup>	21.8	$\pm 1.8$				
	SIG/SIG <sup>odd</sup>	$>\!800$					
	SIG/SIG <sup>even</sup>	57	$\pm 5$				

Table S14. Estimated signal-to-background, background components, and split-initiator HCR suppression. (A) Signal-to-background (SIG/BACK) based on methods of Section S1.4.2. (B) Background components (AF, NSA, NSD) based on methods of Section S1.4.3. (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>) based on methods of Section S1.4.4. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figure S11.

## S3.3 Multiplexed 4-channel mRNA imaging with high signal-to-background in whole-mount chicken embryos (cf. Figure 4)



**Figure S12.** Measurement of signal and background for multiplexed 4-channel mRNA imaging (cf. Figure 4). (A) Individual channels of 4-channel confocal images. For each of three replicate embryos, a representative optical section was selected for each channel based on the expression depth of the corresponding target mRNA. (B) Pixel intensity histograms for Signal + Background (pixels within solid boundary) and Background (pixels within dashed boundary). Ch1: target mRNA *FoxD3*, probe set with 12 split-initiator probe pairs, amplifier B4-Alexa488. Ch2: target mRNA *Dmbx1*, probe set with 20 split-initiator probe pairs, amplifier B1-Alexa514. Ch 3: target mRNA *Sox10*, probe set with 20 split-initiator probe pairs, amplifier B3-Alexa646. Ch4: *EphA4*, probe set with 20 split-initiator probe pairs, amplifier B2-Alexa647. Whole-mount chicken embryos fixed stage HH 10.

\_

Target mRNA	BACK	SIG+BACK	SIG	SIG/BACK
FoxD3 Dmbx1 Sox10 EphA4	$\begin{array}{r} 36 \ \pm 3 \\ 16.4 \pm 0.2 \\ 34.0 \pm 0.2 \\ 35 \ \pm 3 \end{array}$	$\begin{array}{c} 1270 \pm 50 \\ 730 \pm 20 \\ 2030 \pm 80 \\ 980 \pm 70 \end{array}$	$\begin{array}{c} 1230 \pm 50 \\ 710 \pm 20 \\ 2000 \pm 80 \\ 950 \pm 70 \end{array}$	$\begin{array}{rrr} 34 & \pm \ 3 \\ 43.4 \pm 1.0 \\ 59 & \pm \ 3 \\ 27 & \pm \ 3 \end{array}$

Table S15. Estimated signal-to-background for multiplexed 4-channel mRNA imaging. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figure S12 using methods of Section S1.4.2.

## S3.4 qHCR imaging: analog mRNA relative quantitation with subcellular resolution in wholemount chicken embryos (cf. Figure 5)

### S3.4.1 Testing for a crowding effect

In order to perform multiplexed quantitative imaging using HCR, it is important that there is not a crowding effect in which amplification polymers tethered to one target molecule affect the signal intensity for a different target molecule. To test for a possible crowding effect, we imaged two target mRNAs that are highly expressed in the same cells (*EphA4* and *Egr2*) individually (1-target studies) and also simultaneously (2-target studies) within wholemount chicken embryos. Figure S13 compares the signal intensity distributions for 1-target and 2-target studies, revealing similar intensity distributions whether targets were detected alone or together, suggesting that there is not a significant crowding effect (either antagonistic or synergistic).



Figure S13. Comparison of signal intensity distributions for individual and simultaneous imaging of *EphA4* and *Egr2*. (A) Raw voxel intensity scatter plot: *Egr2* channel vs *EphA4* channel. (B) Raw voxel intensity histogram for *EphA4* channel. (C) Raw voxel intensity histogram for *Egr2* channel. In panels B and C, solid lines denote average histograms over 3 replicate embryos while symbols denote individual histograms (1 histogram per replicate). Orange data: signal plus background for *EphA4* and *Egr2* (Figure S14). Cyan data: signal plus background for *EphA4* and background for *EgrA* (Figure S15). Blue data: background for *EphA4* and signal plus background for *Egr2* (Figure S16). Black data (near origin): background for *EphA4* and *Egr2* (Figure S17). Voxel size:  $2.1 \times 2.1 \times 2.7 \mu$ m. Whole-mount wildtype chicken embryos fixed stage HH 10.



Figure S14. Characterizing signal plus background for *EphA4* and *Egr2* in a 2-target experiment. (A) Individual channels from 2-channel confocal images depicting regions used to estimate signal plus background. *EphA4* channel: 20 probe pairs and amplifier B1-Alexa546. *Egr2* channel: 20 probe pairs and amplifier B3-Alexa647. For each of 3 replicate embryos, a representative optical section was selected based on the expression depth of the target mRNAs. Pixel size:  $0.4 \times 0.4 \mu m$ . (B) Raw voxel intensity scatter plots for the selected regions of panel A representing signal plus background for *EphA4* and *Egr2*. (C) Raw voxel intensity histograms for the selected regions of panel A representing signal plus background for *EphA4* and *Egr2*. Same microscope settings used for all replicates in Figures S14–S17. Voxel size:  $2.1 \times 2.1 \times 2.7 \mu m$ . Whole-mount wildtype chicken embryos fixed stage HH 10.



Figure S15. Characterizing signal plus background for *EphA4* in a 1-target experiment. (A) Individual channels from 2-channel confocal images depicting regions used to estimate signal plus background for *EphA4* and background for *Egr2*. *EphA4* channel: 20 probe pairs and amplifier B1-Alexa546. *Egr2* channel: no probes, no amplifier. For each of 3 replicate embryos, a representative optical section was selected based on the expression depth of the target mRNA. Pixel size:  $0.4 \times 0.4$   $\mu$ m. (B) Raw voxel intensity scatter plots for the selected regions of panel A representing signal plus background for *EphA4* and background for *EphA4* and background for *EphA4* and background for *Egr2*. (C) Raw voxel intensity histograms for the selected regions of panel A representing signal plus background for *Egr2*. Same microscope settings used for all replicates in Figures S14–S17. Voxel size:  $2.1 \times 2.1 \times 2.7 \mu$ m. Whole-mount wildtype chicken embryos fixed stage HH 10.



**Figure S16.** Characterizing signal plus background for *Egr2* in a 1-target experiment. (A) Individual channels from 2channel confocal images depicting regions used to estimate background for *EphA4* and signal plus background for *Egr2*. *EphA4* channel: no probes, no amplifier. *Egr2* channel: 20 probe pairs and amplifier B3-Alexa647. For each of 3 replicate embryos, a representative optical section was selected based on the expression depth of the target mRNA. Pixel size:  $0.4 \times 0.4 \,\mu$ m. (B) Raw voxel intensity scatter plots for the selected regions of panel A representing background for *EphA4* and signal plus background for *Egr2*. (C) Raw voxel intensity histograms for the selected regions of panel A representing background for *EphA4* and signal plus background for *Egr2*. Same microscope settings used for all replicates in Figures S14–S17. Voxel size:  $2.1 \times 2.1 \times 2.7 \,\mu$ m. Whole-mount wildtype chicken embryos fixed stage HH 10.



Figure S17. Characterizing background for *EphA4* and *Egr2*. Individual channels from 2-channel confocal images depicting regions used to estimate background using the standard HCR v3.0 in situ protocol (Section S2.1) omitting probes (BACK  $\approx AF + NSA$ ; see Section S1.4 for definitions). For each of 3 replicate embryos, a representative optical section was selected at approximately the depth where *EphA4* and *Egr2* are expressed. Same microscope settings used for all replicates in Figures S14–S17. Pixel size:  $0.4 \times 0.4 \ \mu m$ . (B) Raw voxel intensity scatter plots for the selected region of panel A. (C) Raw voxel intensity histograms for the scatter plots of panel B. Voxel size:  $2.1 \times 2.1 \times 2.7 \ \mu m$ . Whole-mount wildtype chicken embryos fixed stage HH 10.



S3.4.2 Redundant 2-channel detection of Dmbx1

**Figure S18. Redundant 2-channel detection of** *Dmbx1***.** (A) Confocal images: individual channels and merge. Solid boundaries denote regions of high expression; dashed boundaries denote regions of no/low expression. Pixel size:  $0.2 \times 0.2 \mu m$ . Probe sets: 20 split-initiator probe pairs per channel. Amplifiers: B1-Alexa546 (Ch1) and B2-Alexa647 (Ch2). Whole-mount chicken embryos fixed stage HH 10. (B) Raw voxel intensity scatter plots representing signal plus background for voxels within solid boundaries of panel A. Voxel size:  $2.1 \times 2.1 \times 2.7 \mu m$ . Dashed lines represent BOT and TOP values (Table S16) used to normalize data for panel C using methods of Section S1.4.5. (C) Normalized voxel intensity scatter plots representing estimated normalized signal (Pearson correlation coefficient, r).





**Figure S19. Redundant 2-channel detection of** *EphA4* (cf. Figure 5). (A) Confocal images: individual channels and merge. Solid boundaries denote regions of high expression; dashed boundaries denote regions of no/low expression. Pixel size:  $0.2 \times 0.2 \mu m$ . Probe sets: 20 split-initiator probe pairs per channel. Amplifiers: B1-Alexa546 (Ch1) and B2-Alexa647 (Ch2). Whole-mount chicken embryos fixed stage HH 10. (B) Raw voxel intensity scatter plots representing signal plus background for voxels within solid boundaries of panel A. Voxel size:  $2.1 \times 2.1 \times 2.7 \mu m$ . Dashed lines represent BOT and TOP values (Table S16) used to normalize data for panel C using methods of Section S1.4.5. (C) Normalized voxel intensity scatter plots representing estimated normalized signal (Pearson correlation coefficient, r).

Development 145: doi:10.1242/dev.165753: Supplementary information

Target mRNA	Channel	BACK	SIG+BACK	SIG	SIG/BACK	BOT	TOP
Dmbx1 Dmbx1	Alexa546 Alexa647	$\begin{array}{c} 23.9 \pm 1.1 \\ 35 \ \pm 5 \end{array}$	$\begin{array}{c} 840\pm80\\ 770\pm70\end{array}$	$\begin{array}{c} 810\pm80\\ 730\pm70\end{array}$	$\begin{array}{rrr} 34 & \pm  4 \\ 21 & \pm  3 \end{array}$	24 35	3266 3040
EphA4 EphA4	Alexa546 Alexa647	$28.7 \pm 1.1$ $29.7 \pm 1.1$	$1720 \pm 10 \\ 1490 \pm 10$	$1693 \pm 13 \\ 1455 \pm 10$	$59 \pm 2 \\ 48.9 \pm 1.8$	29 30	3995 3855

Table S16. Estimated signal-to-background for redundant 2-channel detection of *Dmbx1* and *EphA4*. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on rectangular regions depicted in Figure S18A and S19A using methods of Section S1.4.2. BOT and TOP values used to calculate normalized voxel intensities for scatter plots of Figures 5C, S18C, and S19C using methods of Section S1.4.5.

## S3.5 In situ validation of automatic background suppression with split-initiator probes for mRNA flow cytometry with cultured human and bacterial cells

The methods of Sections S1.5.2 and S1.5.3 are used to measure:

- signal, background, and signal-to-background (Figures S20A–S22A and Tables S17A–S19A) .
- background components (AF, NSA, NSD; Figures S20A–S22A and Tables S17B–S19B).
- split-initiator HCR suppression (Figures S20B–S22B and Tables S17C–S19C).

Additional measurements of these quantities are provided in the 2-channel experiments of Figures S24–S28 and Tables S20–S24.

S3.5.1 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression for *d2eGFP* transgenic target in HEK cells



**Figure S20.** Measurement of signal and background, background components, and split-initiator HCR suppression for *d2eGFP* transgenic target in HEK cells (cf. Figure 6A). (A) Signal and background: use experiments of Types 1a and 1b in Table S7A (odd + even probes, hairpins) to measure SIG+BACK (GFP+ cells) and BACK (WT cells). Background components: use experiment of Type 2 in Table S7B (no probes, with hairpins) to measure NSA+AF (GFP+ cells); use experiment of Type 3 (no probes, no hairpins) to measure AF (GFP+ cells). (B) Split-initiator HCR suppression: use experiment of Types 4a in Table S7C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (GFP+ cells); use experiment of Type 5a in Table S7C (even probes, hairpins) to measure SIG<sup>even</sup>+BACK<sup>even</sup> (GFP+ cells). Distribution of single-cell fluorescence intensities. Protocol: in situ HCR v3.0 (Section S2.3). Probe set: 12 split-initiator probe pairs. Amplifier: B3-Alexa594. Sample: 55,000 HEK cells in suspension (GFP+ or WT).

		Channel	Reage	ents	
	Quantity	B3-Alexa594	Probes	Hairpins	Cell type
Α	SIG+NSD+NSA+AF = SIG+BACK NSD+NSA+AF = BACK	$\begin{array}{r} 13220 \ \pm 60 \\ 128.5 \pm 0.2 \end{array}$	odd + even odd + even	$\checkmark$	GFP+ WT
	SIG SIG/BACK	$\begin{array}{r} 13090 \ \pm 60 \\ 101.8 \pm 0.5 \end{array}$			
В	NSA+AF AF	$\begin{array}{c} 120.7 \pm 0.5 \\ 104.7 \pm 0.2 \end{array}$		$\checkmark$	GFP+ GFP+
	NSA NSD	$16.0 \pm 0.5$ $7.8 \pm 0.5$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$\begin{array}{c} 149.1 \pm 0.3 \\ 710 \ \pm 3 \end{array}$	odd even	$\checkmark$	GFP+ GFP+
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$\begin{array}{c} 28.4 \pm 0.6 \\ 589 \ \pm 3 \\ 461 \ \pm 9 \\ 22.2 \pm 0.1 \end{array}$			

Table S17. Estimated signal-to-background, background components, and split-initiator HCR suppression for *d2eGFP* transgenic target in HEK cells (cf. Figure 6A). (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA, NSD). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF, which leads to lower bounds on SIG/SIG<sup>odd</sup> and SIG/SIG<sup>even</sup>. Mean  $\pm$  standard error, N = 55,000 cells. Analysis based on single-cell intensities of Figure S20 using methods of Section S1.5.2.

S3.5.2 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression for *GAPDH* endogenous target in HEK cells



**Figure S21. Measurement of signal and background, background components, and split-initiator HCR suppression for** *GAPDH* **endogenous target in HEK cells.** (A) Signal and background: use experiments of Types 1a and 1b in Table S8A to measure SIG+BACK (even + odd probes, hairpins) and BACK (Tg(odd) + Tg(even) probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. (B) Split-initiator HCR suppression: use experiment of Types 4a and 4b in Table S8C to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins) and BACK<sup>odd</sup> (Tg(odd) probes, hairpins); use experiments of Types 5a and 5b in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins) and BACK<sup>even</sup> (Tg(even) probes, hairpins). Distribution of single-cell fluorescence intensities. Protocol: in situ HCR v3.0 (Section S2.3). Probe set: 10 split-initiator probe pairs. Amplifier: B4-Alexa594. Sample: 30,000 HEK cells in suspension (WT).

		Channel	Reagents		
	Quantity	B4-Alexa594	Probes	Hairpins	Cell type
Α	SIG+NSD+NSA+AF = SIG+BACK NSD+NSA+AF = BACK	$\begin{array}{rrr} 4775 & \pm 15 \\ 414.0 & \pm 0.9 \end{array}$	odd + even Tg(odd) + Tg(even)	$\checkmark$	WT WT
	SIG SIG/BACK	$\begin{array}{r} 4362 \\ 10.55 \pm 0.04 \end{array}$			
В	NSA+AF AF	$\begin{array}{rrr} 353.7 & \pm \ 0.8 \\ 304.0 & \pm \ 0.7 \end{array}$		$\checkmark$	WT WT
	NSA NSD	$     50 \pm 1      60 \pm 1 $			
С	SIG <sup>odd</sup> +NSD <sup>odd</sup> +NSA+AF = SIG <sup>odd</sup> +BACK <sup>odd</sup> NSD <sup>odd</sup> +NSA+AF = BACK <sup>odd</sup> SIG <sup>even</sup> +NSD <sup>even</sup> +NSA+AF = SIG <sup>even</sup> +BACK <sup>even</sup> NSD <sup>even</sup> +NSA+AF = BACK <sup>even</sup>	$\begin{array}{rrr} 450 & \pm \ 7 \\ 371.2 & \pm \ 0.8 \\ 499 & \pm \ 1 \\ 397.3 & \pm \ 0.8 \end{array}$	odd Tg(odd) even Tg(even)	✓ ✓ ✓	WT WT WT WT
	NSD <sup>odd</sup> NSD <sup>even</sup> SIG <sup>odd</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$			

Table S18. Estimated signal-to-background, background components, and split-initiator HCR suppression for *GAPDH* endogenous target in HEK cells. (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA, NSD). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). Mean  $\pm$  standard error, N = 30,000 cells. Analysis based on single-cell intensities of Figure S21 using methods of Section S1.5.3.

S3.5.3 Measurement of signal, background, signal-to-background, background components, and split-initiator HCR suppression for *eGFP* transgenic target in *E. coli* 



**Figure S22.** Measurement of signal and background, background components, and split-initiator HCR suppression for *eGFP* transgenic target in *E. coli* (cf. Figure 6A). (A) Signal and background: use experiments of Types 1a and 1b in Table S7A (odd + even probes, hairpins) to measure SIG+BACK (GFP+ cells) and BACK (WT cells). Background components: use experiment of Type 2 in Table S7B (no probes, with hairpins) to measure NSA+AF (GFP+ cells); use experiment of Type 3 (no probes, no hairpins) to measure AF (GFP+ cells). (B) Split-initiator HCR suppression: use experiment of Type 4a in Table S7C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (GFP+ cells); use experiment of Type 5a in Table S7C (even probes, hairpins) to measure SIG<sup>even</sup>+BACK<sup>even</sup> (GFP+ cells). Distribution of single-cell fluorescence intensities. Protocol: in situ HCR v3.0 (Section S2.4). Probe set: 12 split-initiator probe pairs. Amplifier: B3-Alexa594. Sample: 18,000 *E. coli* in suspension (WT: K12 MG1655; GFP+: K12 MG1655 pUA66-sdhC expressing *gfpmut2*).

		Channel	Reage	ents	
	Quantity	B3-Alexa594	Probes	Hairpins	Cell type
Α	SIG+NSD+NSA+AF = SIG+BACK NSD+NSA+AF = BACK	$\begin{array}{rrr} 3330 & \pm 10 \\ 120 & \pm 20 \end{array}$	odd + even odd + even	$\checkmark$	GFP+ WT
	SIG SIG/BACK	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$			
В	NSA+AF AF	$72.5 \pm 0.7 \\ 55.7 \pm 0.7$		$\checkmark$	GFP+ GFP+
	NSA NSD	$\begin{array}{rrr} 17 & \pm 1 \\ 50 & \pm 20 \end{array}$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$\begin{array}{c} 71.3 \pm 0.7 \\ 395 \ \pm 15 \end{array}$	odd even	$\checkmark$	GFP+ GFP+
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup>	$<1 \\ 320 \pm 10 \\ >3000 \\ 20 \pm 0.5 $			
	SIG/SIG <sup>even</sup>	$9.9 \pm 0.5$			

Table S19. Estimated signal-to-background, background components, and split-initiator HCR suppression for *eGFP* transgenic target in *E. coli* (cf. Figure 6A). (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA, NSD). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF, which leads to lower bounds on SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup>. Signal-to-background split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup> and SIG/SIG<sup>odd</sup>.

# S3.6 qHCR flow cytometry: analog mRNA relative quantitation for high-throughput expression profiling of human and bacterial cells (cf. Figure 6)

### S3.6.1 Testing for a crowding effect

In order to perform multiplexed quantitative flow cytometry using HCR, it is important that there is not a crowding effect in which amplification polymers tethered to one target molecule affect the signal intensity for a different target molecule. To test for a possible crowding effect, we analyzed two highly-expressed target mRNAs (*GAPDH* and *ACTB*) individually (1-target studies) and also simultaneously (2-target studies) within HEK cells. Figure S23 compares the signal intensity distributions for 1-target and 2-target studies, revealing similar intensity distributions whether targets were detected alone or together, suggesting that there is not a significant crowding effect (either antagonistic or synergistic).



**Figure S23.** Comparison of signal intensity distributions for individual and multiplexed floHCR of *GAPDH* and *ACTB*. (A) Raw single-cell fluorescence intensity scatter plots: *GAPDH* channel vs *ACTB* channel. (B) Single-cell fluorescence intensity histogram for *GAPDH* channel. (C) Single-cell fluorescence intensity histogram for *ACTB* channel. Orange data: signal plus background for *GAPDH* and *ACTB*. Cyan data: signal plus background for *GAPDH* and autofluorescence for *ACTB*. Blue data: background for *ACTB* and signal plus autofluorescence for *GAPDH*. Black data: autofluorescence for *GAPDH* and *ACTB*. Sample: 65,000 HEK cells in suspension (WT).



#### S3.6.2 Redundant 2-channel detection of GAPDH endogenous target in HEK cells

Figure S24. Measurement of signal and background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *GAPDH* endogenous target (cf. Figure 6B). (A) Signal and background: use experiment of Type 1a in Table S8A to measure SIG+BACK (even + odd probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. Split-initiator HCR suppression: use experiment of Type 4a in Table S8C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins); use experiment of Type 5a in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins). Single-cell fluorescence intensity histograms for Ch1 and Ch2. (B) Raw single-cell fluorescence intensity scatter plots representing signal plus background. Dashed lines represent BOT and TOP values (Table S20) used to normalize data for panel C using methods of Section S1.5.4 (outliers excluded from normalized scatter plots marked with circles). (C) Normalized single-cell fluorescence intensity scatter plots representing estimated normalized signal (Pearson correlation coefficient, *r*). Protocol: in situ HCR v3.0 (Section S2.3). Probe sets: 10 split-initiator probe pairs per channel. Amplifiers: B5-Alexa488 (Ch1) and B4-Alexa594 (Ch2). Sample: 20,000 HEK cells in suspension (WT).

		Cha	nnel	Reage	ents	
	Quantity	Ch1: B5-Alexa488	Ch2: B4-Alexa594	Probes	Hairpins	Cell type
Α	SIG+BACK	$870 \pm 5$	$3105 \pm 14$	odd + even	$\checkmark$	WT
	SIG SIG/BACK	$808 \pm 5 \\ 12.86 \pm 0.08$	$\begin{array}{r} 2904 \\ 14.43 \pm 0.08 \end{array}$			
В	NSA+AF AF	$\begin{array}{rrr} 62.8 & \pm \ 0.2 \\ 57.4 & \pm \ 0.2 \end{array}$	$\begin{array}{rrr} 201.2 & \pm \ 0.6 \\ 166.5 & \pm \ 0.5 \end{array}$		$\checkmark$	WT WT
	NSA	$5.4 \pm 0.3$	$34.7 \pm 0.8$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$\begin{array}{rrr} 82.8 & \pm \ 0.3 \\ 63.0 & \pm \ 0.2 \end{array}$	$\begin{array}{rr} 244.5 & \pm \ 0.8 \\ 256.6 & \pm \ 0.8 \end{array}$	odd even	$\checkmark$	WT WT
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$\begin{array}{rrr} 20.0 & \pm \ 0.4 \\ < 0.3 \\ 40.4 & \pm \ 0.8 \\ > 3000 \end{array}$	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$			
D	BOT TOP	$62.8 \\ 5265.2$	$\begin{array}{c} 201.2 \\ 19049.4 \end{array}$			

Table S20. Estimated signal-to-background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *GAPDH* endogenous target (cf. Figure 6B). (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG, SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK = BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF. Mean  $\pm$ standard error, N = 20,000 cells. Analysis based on single-cell intensities of Figure S24 using methods of Section S1.5.3. (D) BOT and TOP values used to calculate normalized single-cell intensities for scatter plots of Figures 6B and S24C using methods of Section S1.5.4.



#### S3.6.3 Redundant 2-channel detection of PGK1 endogenous target in HEK cells

Figure S25. Measurement of signal and background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *PGK1* endogenous target. (A) Signal and background: use experiment of Type 1a in Table S8A to measure SIG+BACK (even + odd probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. Split-initiator HCR suppression: use experiment of Type 4a in Table S8C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins); use experiment of Type 5a in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins). Singlecell fluorescence intensity histograms for Ch1 and Ch2. (B) Raw single-cell fluorescence intensity scatter plots representing signal plus background. Dashed lines represent BOT and TOP values (Table S21) used to normalize data for panel C using methods of of Section S1.5.4 (outliers excluded from normalized scatter plots marked with circles). (C) Normalized single-cell fluorescence intensity scatter plots representing estimated normalized signal (Pearson correlation coefficient, *r*). Protocol: in situ HCR v3.0 (Section S2.3). Probe sets: 18 split-initiator probe pairs per channel. Amplifiers: B1-Alexa488 (Ch1) and B2-Alexa594 (Ch2). Sample: 54,000 HEK cells in suspension (WT).
		Ch	annel	Reage	Reagents	
	Quantity	Ch1: B1-Alexa488	Ch2: B2-Alexa594	Probes	Hairpins	Cell type
Α	SIG+BACK	$4145 \pm 11$	$1528 \pm 4$	odd + even	$\checkmark$	WT
	SIG SIG/BACK	$\begin{array}{c} 3843 \\ 12.72 \pm 0.04 \end{array}$	$\begin{array}{c} 1248 \\ 4.47 \pm 0.02 \end{array}$			
В	NSA+AF AF	$\begin{array}{rrr} 302.1 & \pm \ 0.5 \\ 289.3 & \pm \ 0.5 \end{array}$	$\begin{array}{rr} 279.4 & \pm \ 0.5 \\ 220.1 & \pm \ 0.4 \end{array}$		$\checkmark$	WT WT
	NSA	$12.9 \hspace{0.2cm} \pm \hspace{0.2cm} 0.7$	$59.3 \pm 0.6$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$\begin{array}{rrr} 301.3 & \pm \ 0.5 \\ 380.5 & \pm \ 0.6 \end{array}$	$\begin{array}{ccc} 309.5 & \pm  0.6 \\ 374 & \pm  7 \end{array}$	odd even	$\checkmark$	WT WT
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	< 0.7 78.4 $\pm 0.8$ >5000 49.0 $\pm 0.5$	$\begin{array}{rrrr} 30.1 & \pm 0.8 \\ 94 & \pm 7 \\ 41 & \pm 1 \\ 13 & \pm 1 \end{array}$			
D	BOT TOP	$302.1 \\ 35538.8$	279.4 11 847.7			

Table S21. Estimated signal-to-background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *PGK1* endogenous target. (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG, SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK = BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF. Mean  $\pm$  standard error, N = 54,000 cells. Analysis based on single-cell intensities of Figure S25 using methods of Section S1.5.3. (D) BOT and TOP values used to calculate normalized single-cell intensities for scatter plots of Figure S25C using methods of Section S1.5.4.



#### S3.6.4 Redundant 2-channel detection of *fusA* endogenous target in *E. coli*

Figure S26. Measurement of signal and background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *fusA* endogenous target (cf. Figure 6B). (A) Signal and background: use experiment of Type 1a in Table S8A to measure SIG+BACK (even + odd probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. Split-initiator HCR suppression: use experiment of Type 4a in Table S8C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins); use experiment of Type 5a in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins). Single-cell fluorescence intensity histograms for Ch1 and Ch2. (B) Raw single-cell fluorescence intensity scatter plots representing signal plus background. Dashed lines represent BOT and TOP values (Table S22) used to normalize data for panel C using methods of of Section S1.5.4 (outliers excluded from normalized scatter plots marked with ellipses). (C) Normalized single-cell fluorescence intensity scatter plots representing estimated normalized signal (Pearson correlation coefficient, *r*). Protocol: in situ HCR v3.0 (Section S2.4). Probe sets: 18 split-initiator probe pairs per channel. Amplifiers: B3-Alexa488 (Ch1) and B2-Alexa594 (Ch2). Sample: 3,400 *E. coli* K12 MG1655 in suspension (WT).

		Ch	annel	Reage	ents	
	Quantity	Ch1: B3-Alexa488	Ch2: B2-Alexa594	Probes	Hairpins	Cell type
Α	SIG+BACK	$13100\pm300$	$15800\pm300$	odd + even	$\checkmark$	WT
	SIG SIG/BACK	$\begin{array}{rrr} 13000 & \pm 300 \\ 99 & \pm 3 \end{array}$	$\begin{array}{rrr} 15700 & \pm 300 \\ 135 & \pm 7 \end{array}$			
В	NSA+AF AF	$\begin{array}{rrr} 130 & \pm \ 3 \\ 126 & \pm \ 7 \end{array}$	$\begin{array}{rrr} 116 & \pm \ 5 \\ 120 & \pm \ 10 \end{array}$		$\checkmark$	WT WT
	NSA	<7	<14			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$500 \pm 100 \\ 1100 \pm 200$	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	odd even	$\checkmark$	WT WT
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$\begin{array}{rrrr} 300 & \pm 100 \\ 900 & \pm 200 \\ 40 & \pm 20 \\ 14 & \pm 3 \end{array}$	$\begin{array}{rrr} 300 & \pm 100 \\ 2500 & \pm 200 \\ 50 & \pm 20 \\ 6.2 \pm 0.6 \end{array}$			
D	BOT TOP	$\frac{130.3}{150951.6}$	$\frac{116}{183947.3}$			

Table S22. Estimated signal-to-background, background components, and split-initiator HCR suppression for redundant 2-channel detection of *fusA* endogenous target (cf. Figure 6B). (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG, SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK = BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF. Mean  $\pm$ standard error, N = 3,400 cells. Analysis based on single-cell intensities of Figure S26 using methods of Section S1.5.3. (D) BOT and TOP values used to calculate normalized single-cell intensities for scatter plots of Figures 6B and S26C using methods of Section S1.5.4.



#### S3.6.5 Multiplexed 2-channel detection of GAPDH and PGK1 endogenous targets in HEK cells

**Figure S27.** Measurement of signal and background, background components, and split-initiator HCR suppression for multiplexed 2-channel detection of *GAPDH* and *PGK1* endogenous targets. (A) Single-cell fluorescence intensity histograms for Ch1 and Ch2. (B) Raw single-cell fluorescence intensity scatter plots for Ch1 vs Ch2. Signal and background: use experiment of Type 1a in Table S8A to measure SIG+BACK (even + odd probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. (C) Split-initiator HCR suppression: use experiment of Type 4a in Table S8C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins); use experiment of Type 5a in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins). Protocol: in situ HCR v3.0 (Section S2.3). Ch1: target mRNA *GAPDH*, probe set with 10 splitinitiator probe pairs, amplifier B4-Alexa488. Ch2: target mRNA *PGK1*, probe set with 18 split-initiator probe pairs, amplifier B2-Alexa594. Sample: 18,000 HEK cells in suspension (WT).

		Cha	nnel	Reage	ents	
	Quantity	Ch1: B4-Alexa488	Ch2: B2-Alexa594	Probes	Hairpins	Cell type
Α	SIG+BACK	$6980 \pm 30$	$2073 \pm 8$	odd + even	$\checkmark$	WT
	SIG SIG/BACK	$\begin{array}{cc} 6760 \pm 30 \\ 29.6 \pm 0.1 \end{array}$	$\begin{array}{r} 1806 \\ 6.77 \pm 0.05 \end{array}$			
В	NSA+AF AF	$228.2 \pm 0.7$ $219.7 \pm 0.6$	$\begin{array}{rr} 266.6 & \pm  1.5 \\ 198.4 & \pm  0.6 \end{array}$		$\checkmark$	WT WT
	NSA	$8.4\pm0.9$	$68 \pm 2$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	$355 \pm 9 \\ 362.8 \pm 1.2$	odd even	$\checkmark$	WT WT
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$\begin{array}{rrrr} 73 & \pm 5 \\ 74 & \pm 14 \\ 93 & \pm 6 \\ 91 & \pm 14 \end{array}$	$\begin{array}{rrr} 88 & \pm 9 \\ 96 & \pm 2 \\ 21 & \pm 2 \\ 18.8 & \pm 0.4 \end{array}$			

Table S23. Estimated signal-to-background, background components, and split-initiator HCR suppression for multiplexed 2-channel detection of *GAPDH* and *PGK1* endogenous targets. (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG, SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK = BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF. Mean  $\pm$ standard error, N = 18,000 cells. Analysis based on single-cell intensities of Figure S27 using methods of Section S1.5.3.





**Figure S28.** Measurement of signal and background, background components, and split-initiator HCR suppression for multiplexed 2-channel detection of *fusA* and *icd* endogenous targets. (A) Single-cell fluorescence intensity histograms for Ch1 and Ch2. (B) Raw single-cell fluorescence intensity scatter plots for Ch1 vs Ch2. Signal and background: use experiment of Type 1a in Table S8A to measure SIG+BACK (even + odd probes, hairpins). Background components: use experiment of Type 2 in Table S8B (no probes, with hairpins) to measure NSA+AF; use experiment of Type 3 (no probes, no hairpins) to measure AF. (C) Split-initiator HCR suppression: use experiment of Type 4a in Table S8C (odd probes, hairpins) to measure SIG<sup>odd</sup>+BACK<sup>odd</sup> (odd probes, hairpins); use experiment of Type 5a in Table S8C to measure SIG<sup>even</sup>+BACK<sup>even</sup> (even probes, hairpins). Protocol: in situ HCR v3.0 (Section S2.4). Ch1: target mRNA *fusA*, probe set with 18 split-initiator probe pairs, amplifier B3-Alexa488. Ch2: target mRNA *icd*, probe set with 20 split-initiator probe pairs, amplifier B1-Alexa594. Sample: 35,000 *E. coli* K12 MG1655 in suspension (WT).

		Cha	nnel	Reage	ents	
	Quantity	Ch1: B3-Alexa488	Ch2: B1-Alexa594	Probes	Hairpins	Cell type
Α	SIG+BACK	$1756 \pm 9$	$2533 \pm 12$	odd + even	$\checkmark$	WT
	SIG SIG/BACK	$egin{array}{ccc} 1673 & \pm \ 9 \\ 20.1 \pm 0.2 \end{array}$	$\begin{array}{rrr} 2470 & \pm 12 \\ 38.9 & \pm 0.5 \end{array}$			
В	NSA+AF AF	$83.2 \pm 0.7$ $82.3 \pm 0.7$	$\begin{array}{rrr} 63.5 & \pm \ 0.7 \\ 60.1 & \pm \ 0.7 \end{array}$		$\checkmark$	WT WT
	NSA	$1 \pm 1$	$3.4\pm0.9$			
С	SIG <sup>odd</sup> +BACK <sup>odd</sup> SIG <sup>even</sup> +BACK <sup>even</sup>	$     \begin{array}{r}       86 & \pm 1 \\       180 & \pm 8     \end{array} $	$\begin{array}{ccc} 66.6 & \pm \ 0.8 \\ 93 & \pm \ 1 \end{array}$	odd even	$\checkmark$	WT WT
	SIG <sup>odd</sup> SIG <sup>even</sup> SIG/SIG <sup>odd</sup> SIG/SIG <sup>even</sup>	$ \begin{array}{r} 3  \pm 1 \\ 97  \pm 8 \\ 600  \pm 300 \\ 17.2 \pm 1.3 \end{array} $	$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$			

Table S24. Estimated signal-to-background, background components, and split-initiator HCR suppression for multiplexed 2-channel detection of *fusA* and *icd* endogenous targets. (A) Signal-to-background (SIG/BACK). (B) Background components (AF, NSA). (C) Split-initiator HCR suppression (SIG/SIG<sup>odd</sup>, SIG/SIG<sup>even</sup>). The signal estimates SIG, SIG<sup>odd</sup> and SIG<sup>even</sup> are calculated using the background approximation BACK = BACK<sup>odd</sup> = BACK<sup>even</sup>  $\approx$  NSA+AF. Mean  $\pm$  standard error, N = 35,000 cells. Analysis based on single-cell intensities of Figure S28 using methods of Section S1.5.3.





Figure S29. Redundant 2-channel detection of single *BRAF* mRNAs in HEK cells using in situ HCR v3.0 (cf. Figure 7A). Confocal images: individual channels and merge (without and with DAPI nuclear stain). Maximum intensity projection in the axial direction over 7.14  $\mu$ m (17 focal planes). Pixel size: 0.062×0.062  $\mu$ m. Probe sets: 23 split-initiator probe pairs per channel. Amplifiers: B3-Alexa647 (Ch1) and B4-Alexa546 (Ch2). Red circles: dots detected in Ch1. Green circles: dots detected in Ch2. Yellow circles: dots detected in both channels.

	Dots		Colocalized dots	Colocalizatio	on fractions
	$N_1$ $N_2$		N12	$C_1$	$C_2$
Replicate 1	129	136	110	0.85	0.81
Replicate 2	63	65	53	0.84	0.82
Replicate 3	170	170	144	0.85	0.85
Mean			$0.85\pm0.003$	$0.82\pm0.01$	

Table S25. Dot colocalization fractions for redundant 2-channel detection of single *BRAF* mRNAs in HEK cells using in situ HCR v3.0 (cf. Figure 7A). Mean  $\pm$  standard error, N = 3 replicate samples. Analysis based on the images of Figure S29 using the methods of Section S1.4.6 with the settings in Table S6.

# S3.7.2 Redundant 2-channel detection of single *Dmbx1* mRNAs in whole-mount chicken embryos using in situ HCR v3.0



Figure S30. Redundant 2-channel detection of single *Dmbx1* mRNAs in whole-mount chicken embryos using in situ HCR v3.0 (cf. Figure 7B). Confocal images: individual channels and merge. Maximum intensity projection in the axial direction over 5.04-23.52  $\mu$ m (12, 54, 56 focal planes for replicates 1, 2, 3 depending on sample thickness). Pixel size: 0.099×0.099  $\mu$ m. Probe sets: 25 split-initiator probe pairs per channel. Amplifiers: B2-Alexa647 (Ch1) and B1-Alexa594 (Ch2). Embryos fixed stage HH 8. Red circles: dots detected in Ch1. Green circles: dots detected in Ch2. Yellow circles: dots detected in both channels.

	Dots		Colocalized dots	Colocalizati	tion fractions		
	$N_1$	$N_2$	N <sub>12</sub>	$C_1$	$C_2$		
Replicate 1	403	417	364	0.90	0.87		
Replicate 2	992	990	794	0.80	0.80		
Replicate 3	526	539	448	0.85	0.83		
Mean				$0.85\pm0.03$	$0.84\pm0.02$		

Table S26. Dot colocalization fractions for redundant 2-channel detection of single Dmbx1 mRNAs in whole-mount chicken embryos using in situ HCR v3.0 (cf. Figure 7B). Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on the images of Figure S30 using the methods of Section S1.4.6 with the settings in Table S6.

# S3.7.3 Redundant 2-channel detection of single *kdrl* mRNAs in whole-mount zebrafish embryos using in situ HCR v2.0 (Shah *et al.*, 2016)



Figure S31. Redundant 2-channel detection of single *kdrl* mRNAs in whole-mount zebrafish embryos using in situ HCR v2.0. Spinning disk confocal images: individual channels and merge from Shah *et al.* (2016). Maximum intensity projection in the axial direction over 13  $\mu$ m (39 focal planes). Pixel size: 0.217×0.217  $\mu$ m. Probe sets: 39 standard probes per channel, each incorporating a 30-nt target-binding domain and a full HCR initiator. Amplifiers: B3-Alexa647 (Ch1) and B2-Alexa546 (Ch2). Embryos fixed 27 hpf. Red circles: dots detected in Ch1. Green circles: dots detected in Ch2. Yellow circles: dots detected in both channels.

	Dots		Colocalized dots	Colocalization fraction	
	$N_1$ $N_2$		N <sub>12</sub>	$C_1$	$C_2$
Replicate 1	139	132	79	0.57	0.60
Replicate 2	220	215	113	0.51	0.53
Replicate 3	243	245	91	0.37	0.37
Mean			$0.49\pm0.06$	$0.50\pm0.07$	

Table S27. Dot colocalization fractions for redundant 2-channel detection of single *kdrl* mRNAs in whole-mount zebrafish embryos using in situ HCR v3.0. Mean  $\pm$  standard error, N = 3 replicate embryos. Analysis based on the images of Figure S31 using the methods of Section S1.4.6 with the settings in Table S6.

## S4 Probe sequences

Target mRNA sequences were obtained from the National Center for Biotechnology Information (NCBI) (McEntyre & Ostell, 2002) with the exception of d2eGFP (pd2EGFP-1, Clontech, Cat. #6008-1). Spatial and temporal expression information for whole-mount chicken embryos were obtained from the Gallus Expression in Situ Hybridization Analysis (GEISHA) (Bell *et al.*, 2004; Darnell *et al.*, 2007). Within a given probe set, each DNA standard probe or split-initiator probe pair initiates the same DNA HCR amplifier. Sequences are listed 5' to 3'. Probes are numbered consecutively moving along a target mRNA. For redundant detection experiments, two probe sets are used with each probe set taking alternating probe pairs from along the target (this leads to non-consecutive numbers within each probe set).

## S4.1 Standard probes for Figures 3, S5, and S7

Organism: G. gallus domesticus

Target mRNA: **SRY** (**sex determining region Y**)-**box 10** (*Sox10*) Probe set: **5, 10, or 20 probes (each carrying 2 HCR initiators)** HCR amplifier: **B3-Alexa647** 

#	Initiator I1	Spacer	Probe Sequence (50 nt)	Spacer	Initiator I2
1	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CATggACCCgTCACTCCATgTCTTgAgTCTTCCTCATCTAgAAggCCAAT	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
2	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCAgCAgggATCAAgATTCATgCATgTgTgAATCTTAggCAggACTgCTg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
3	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CgggCTATgAAATgAgAAAggCTAAggCTgACAgTgCAgTTCCTgAATCC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
4	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TTCACgTTTTCAgCAgACACAgTCAAATgCTggAggAgCAAggACCTggT	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
5	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	$\verb ATTggAACCACATCTgggTgTTggCAAgTgCATggTAgCTTTCTTggTgC  $	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
6	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AgAggCggggAgAAAAgCTATAgCgTgCAgCTgTgAAAATCAgCAAggAA	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
7	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	ATAAAATCCATgCAggAAggggTgTgggATTAAACAgATgggACAggggg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
8	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gATggCgATAATgTgATgAACAAACgAgCAgTgATgTACACCCCATCggC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
9	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	ATCCACgAgAgTATCTTTCCATCCTgAgTgAAAgTAggAgggAg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
10	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	ACCCgTTAgAAggTCCCACAACACATCTCTCTgATCAgTTgTCAggAgTC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
11	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCggCgAgAggCAgTggTggTCTTCAgAACCCACTgggCTCATCTCCACC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
12	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCTCgCCCTgCgTggCCTTgCCATTTTTCCgCCTCCgTggCTggTACTTg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
13	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TTCTTgTAgTgAgCCTggATAgAggCAgCCCCgCCAgCCTCCCCCCCAC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
14	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TgCCCATCggACATgggTgACCCTTCCCCAggATgCCTgTggTCCAggTg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
15	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TTAggggTggTgggAggAgTgggAgggCCgTggCTCTgACCTgAAgAgTg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
16	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCCCAAgggAACgCCCTTCTCgCTTggAgTCAgCTTTgCCTgCC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
17	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	ggCCTgggTggCCggCgTgTCCgTTgggTggCAggTATTggTCAAATTCg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
18	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TCCATgCTgCTTggAgATCCAggCTgAgTgTCCACTggCCgCAgCCAggg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
19	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AACCCTgTgAAggCTgCAgCTCCTgTCCCAgTgCCCCCCTgTTCTCCCCTC	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC
20	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gggTCCAgTCATAgCCgCTCAgCACCTggCTgACCgCTTCACggATgCAg	TAAAA	AAAgTCTAATCCgTCCCTgCCTCTATATCTCCACTC

## S4.2 Split-initiator probes for Figures 3, S6, S8, S9, and S10

Organism: G. gallus domesticus

Target mRNA: SRY (sex determining region Y)-box 10 (Sox10)

#### Probe set: 5, 10, or 20 split-initiator probe pairs (each probe carries half an HCR initiator)

HCR amplifier: B3-Alexa647

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	gTCTTCCTCATCTAgAAggCCAATA	2	CCATggACCCgTCACTCCATgTCTT	TT	CCACTCAACTTTAACCCg
3	gTCCCTgCCTCTATATCT	TT	TgTgAATCTTAggCAggACTgCTgC	4	TCCAgCAgggATCAAgATTCATgCA	TT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	TT	gCTgACAgTgCAgTTCCTgAATCCT	6	ACgggCTATgAAATgAgAAAggCTA	TT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	TT	TgCTggAggAgCAAggACCTggTCT	8	TTCACgTTTTCAgCAgACACAgTCA	TT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	TT	CAAgTgCATggTAgCTTTCTTggTg	10	AATATTggAACCACATCTgggTgTT	TT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	TT	AgCTgTgAAAATCAgCAAggAAgCA	12	gAggCggggAgAAAAgCTATAgCgT	TT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	TT	gggATTAAACAgATgggACAggggg	14	TTATAAAATCCATgCAggAAggggT	TT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	ΤT	AgCAgTgATgTACACCCCATCggCC	16	AgATggCgATAATgTgATgAACAAA	TT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	TT	TgAgTgAAAgTAggAgggAggTgCT	18	TATATCCACgAgAgTATCTTTCCAT	TT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	CTCTCTgATCAgTTgTCAggAgTCA	20	TACCCgTTAgAAggTCCCACAACAC	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	gAACCCACTgggCTCATCTCCACCT	22	CCCggCgAgAggCAgTggTggTCTT	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	ΤT	TTCCgCCTCCgTggCTggTACTTgT	24	CCCTCgCCCTgCgTggCCTTgCCAT	TT	CCACTCAACTTTAACCCg
25	gTCCCTgCCTCTATATCT	TT	AgCCCCgCCAgCCTCCCCCTCCACC	26	ATTCTTgTAgTgAgCCTggATAgAg	TT	CCACTCAACTTTAACCCg
27	gTCCCTgCCTCTATATCT	ΤT	CCCAggATgCCTgTggTCCAggTgg	28	gTgCCCATCggACATgggTgACCCT	TT	CCACTCAACTTTAACCCg
29	gTCCCTgCCTCTATATCT	ΤT	gCCgTggCTCTgACCTgAAgAgTgC	30	CTTAggggTggTgggAggAgTgggA	TT	CCACTCAACTTTAACCCg
31	gTCCCTgCCTCTATATCT	ΤT	gAgTCAgCTTTgCCTgCCTgCAgCT	32	TCCCCAAgggAACgCCCTTCTCgCT	TT	CCACTCAACTTTAACCCg
33	gTCCCTgCCTCTATATCT	ΤT	ggTggCAggTATTggTCAAATTCgT	34	TggCCTgggTggCCggCgTgTCCgT	TT	CCACTCAACTTTAACCCg
35	gTCCCTgCCTCTATATCT	ΤT	AgTgTCCACTggCCgCAgCCAgggC	36	CTCCATgCTgCTTggAgATCCAggC	TT	CCACTCAACTTTAACCCg
37	gTCCCTgCCTCTATATCT	ΤT	CCCAgTgCCCCCCTgTTCTCCCTCC	38	AAACCCTgTgAAggCTgCAgCTCCT	TT	CCACTCAACTTTAACCCg
39	gTCCCTgCCTCTATATCT	ΤT	TggCTgACCgCTTCACggATgCAgA	40	AgggTCCAgTCATAgCCgCTCAgCA	TT	CCACTCAACTTTAACCCg

## S4.3 Standard probes for Figures S9 and S10

Organism: *G. gallus domesticus* Target mRNA: **SRY (sex determining region Y)-box 10** (*Sox10*) Probe set: **20 probe pairs (odd probes carry full HCR initiator, even probes carry no initiator)** HCR amplifier: **B3-Alexa647** 

Odd #	Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)
1	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gTCTTCCTCATCTAgAAggCCAATA	2	CCATggACCCgTCACTCCATgTCTT
3	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TgTgAATCTTAggCAggACTgCTgC	4	TCCAgCAgggATCAAgATTCATgCA
5	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gCTgACAgTgCAgTTCCTgAATCCT	6	ACgggCTATgAAATgAgAAAggCTA
7	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TgCTggAggAgCAAggACCTggTCT	8	TTCACgTTTTCAgCAgACACAgTCA
9	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CAAgTgCATggTAgCTTTCTTggTg	10	AATATTggAACCACATCTgggTgTT
11	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AgCTgTgAAAATCAgCAAggAAgCA	12	gAggCggggAgAAAAgCTATAgCgT
13	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gggATTAAACAgATgggACAggggg	14	TTATAAAATCCATgCAggAAggggT
15	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AgCAgTgATgTACACCCCATCggCC	16	AgATggCgATAATgTgATgAACAAA
17	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TgAgTgAAAgTAggAgggAggTgCT	18	TATATCCACgAgAgTATCTTTCCAT
19	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CTCTCTgATCAgTTgTCAggAgTCA	20	TACCCgTTAgAAggTCCCACAACAC
21	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gAACCCACTgggCTCATCTCCACCT	22	CCCggCgAgAggCAgTggTggTCTT
23	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TTCCgCCTCCgTggCTggTACTTgT	24	CCCTCgCCCTgCgTggCCTTgCCAT
25	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AgCCCCgCCAgCCTCCCCCTCCACC	26	ATTCTTgTAgTgAgCCTggATAgAg
27	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCCAggATgCCTgTggTCCAggTgg	28	gTgCCCATCggACATgggTgACCCT
29	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gCCgTggCTCTgACCTgAAgAgTgC	30	CTTAggggTggTgggAggAgTgggA
31	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	gAgTCAgCTTTgCCTgCCTgCAgCT	32	TCCCCAAgggAACgCCCTTCTCgCT
33	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	ggTggCAggTATTggTCAAATTCgT	34	TggCCTgggTggCCggCgTgTCCgT
35	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	AgTgTCCACTggCCgCAgCCAgggC	36	CTCCATgCTgCTTggAgATCCAggC
37	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	CCCAgTgCCCCCCTgTTCTCCCTCC	38	AAACCCTgTgAAggCTgCAgCTCCT
39	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg	TACAA	TggCTgACCgCTTCACggATgCAgA	40	AgggTCCAgTCATAgCCgCTCAgCA

#### Organism: *G. gallus domesticus* Target mRNA: **SRY (sex determining region Y)-box 10** (*Sox10*) Probe set: **20 probe pairs (odd probes carry no initiator, even probes carry full HCR initiator)** HCR amplifier: **B3-Alexa647**

Odd #	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	Initiator I1
1	gTCTTCCTCATCTAgAAggCCAATA	2	CCATggACCCgTCACTCCATgTCTT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
3	TgTgAATCTTAggCAggACTgCTgC	4	TCCAgCAgggATCAAgATTCATgCA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
5	gCTgACAgTgCAgTTCCTgAATCCT	6	ACgggCTATgAAATgAgAAAggCTA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
7	TgCTggAggAgCAAggACCTggTCT	8	TTCACgTTTTCAgCAgACACAgTCA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
9	CAAgTgCATggTAgCTTTCTTggTg	10	AATATTggAACCACATCTgggTgTT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
11	AgCTgTgAAAATCAgCAAggAAgCA	12	gAggCggggAgAAAAgCTATAgCgT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
13	gggATTAAACAgATgggACAggggg	14	TTATAAAATCCATgCAggAAggggT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
15	AgCAgTgATgTACACCCCATCggCC	16	AgATggCgATAATgTgATgAACAAA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
17	TgAgTgAAAgTAggAgggAggTgCT	18	TATATCCACgAgAgTATCTTTCCAT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
19	CTCTCTgATCAgTTgTCAggAgTCA	20	TACCCgTTAgAAggTCCCACAACAC	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
21	gAACCCACTgggCTCATCTCCACCT	22	CCCggCgAgAggCAgTggTggTCTT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
23	TTCCgCCTCCgTggCTggTACTTgT	24	CCCTCgCCCTgCgTggCCTTgCCAT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
25	AgCCCCgCCAgCCTCCCCCTCCACC	26	ATTCTTgTAgTgAgCCTggATAgAg	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
27	CCCAggATgCCTgTggTCCAggTgg	28	gTgCCCATCggACATgggTgACCCT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
29	gCCgTggCTCTgACCTgAAgAgTgC	30	CTTAggggTggTgggAggAgTgggA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
31	gAgTCAgCTTTgCCTgCCTgCAgCT	32	TCCCCAAgggAACgCCCTTCTCgCT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
33	ggTggCAggTATTggTCAAATTCgT	34	TggCCTgggTggCCggCgTgTCCgT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
35	AgTgTCCACTggCCgCAgCCAgggC	36	CTCCATgCTgCTTggAgATCCAggC	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
37	CCCAgTgCCCCCCTgTTCTCCCTCC	38	AAACCCTgTgAAggCTgCAgCTCCT	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg
39	TggCTgACCgCTTCACggATgCAgA	40	AgggTCCAgTCATAgCCgCTCAgCA	TACTT	gTCCCTgCCTCTATATCTCCACTCAACTTTAACCCg

## S4.4 Split-initiator probes for Figure S11

Organism: G. gallus domesticus

#### Target mRNA: EPH receptor A4 (*EphA4*)

#### Probe set: 20 split-initiator probe pairs (each probe carries half an HCR initiator)

HCR amplifier: **B2-Alexa647** 

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCgTAAATCCTCATCA	AA	gCgCCgACggggACCCCggCCATgC	2	CAgACgCCgACgAggAgCgggAggA	AA	ATCATCCAgTAAACCgCC
5	CCTCgTAAATCCTCATCA	AA	CCCAgCTCCCCCTgCACCgAgCggg	6	CCTCCTTCCAgCgggCTCgCgATCC	AA	ATCATCCAgTAAACCgCC
9	CCTCgTAAATCCTCATCA	AA	TgACTgggCTCCATCACATTgCAAA	10	ATCCAATCAgTTCgTAACCAATTAT	AA	ATCATCCAgTAAACCgCC
13	CCTCgTAAATCCTCATCA	AA	TCCTTgTCgTTgTTTgATTCATAgT	14	gCAAACTggCTCTCTCgAATAAAAC	AA	ATCATCCAgTAAACCgCC
17	CCTCgTAAATCCTCATCA	AA	AgTgggCACTTCTTATAgAAgACAC	18	ggAAACTgTgCCAggTTTCgAACTg	AA	ATCATCCAgTAAACCgCC
21	CCTCgTAAATCCTCATCA	AA	ACgTCCTTCTCTTCCgAgTTgTTgA	22	CCATCTgCCCCgCAgTACATTTTTg	AA	ATCATCCAgTAAACCgCC
25	CCTCgTAAATCCTCATCA	AA	ATTTTgCAAgCTTggCATTCACCAT	26	TCTgTTgAgAgCgCCTTgTAgTATC	AA	ATCATCCAgTAAACCgCC
29	CCTCgTAAATCCTCATCA	AA	AAgAAgCCCCgATCACAggTgCAgg	30	ATggATgCAgCATCATTTTCTgCTC	AA	ATCATCCAgTAAACCgCC
33	CCTCgTAAATCCTCATCA	AA	gCgCTCCACTCCAAgTTCACTgACg	34	TCgTCCCgTCCTCCCTTgTTCTgTg	AA	ATCATCCAgTAAACCgCC
37	CCTCgTAAATCCTCATCA	AA	AAATgTACACCACTgCCACAggACC	38	gTTTTCAgCCCgTTCTgCTgggggC	AA	ATCATCCAgTAAACCgCC
41	CCTCgTAAATCCTCATCA	AA	TgCTTggACACTCCATTCACTgCCC	42	gACACAgCTTggTCCTggCTggggT	AA	ATCATCCAgTAAACCgCC
45	CCTCgTAAATCCTCATCA	AA	gCAACgCTgTgCCTCgTTATCTCTT	46	ggCCTgTCAggTTCCAgCCAggCCA	AA	ATCATCCAgTAAACCgCC
49	CCTCgTAAATCCTCATCA	AA	gCTgTCTTCACAATgCgATAgCTgC	50	AAACCTTTgATgTCAgTATTCCTgg	AA	ATCATCCAgTAAACCgCC
53	CCTCgTAAATCCTCATCA	AA	AACTCAAACggCCCACTgAAgTCTC	54	ATgggggAAggAACTgTgTTAgTTg	AA	ATCATCCAgTAAACCgCC
57	CCTCgTAAATCCTCATCA	AA	gCTgCAATgAgAATgACCACgAgAA	58	TTgCTgCgCCTCCTgCTgATgACAA	AA	ATCATCCAgTAAACCgCC
61	CCTCgTAAATCCTCATCA	AA	TATGTAAAAggATCCACATATGTTC	62	TCCCTCACAgCTTgATTTggATCCT	AA	ATCATCCAgTAAACCgCC
65	CCTCgTAAATCCTCATCA	AA	CATTTAgTAACAACgCCTTCCAAgT	66	TACTCAgTTATgATCATTACTggTT	AA	ATCATCCAgTAAACCgCC
69	CCTCgTAAATCCTCATCA	AA	TCCCgATgCACATAgCTCATgTCAg	70	TTgACCAgTATgTTTCgAgCAgCTA	AA	ATCATCCAgTAAACCgCC
73	CCTCgTAAATCCTCATCA	AA	CCAAATTTAggTCTgTCgCTgCgTT	74	AgTTTgTCCAgCATgTTgACAATCT	AA	ATCATCCAgTAAACCgCC
77	CCTCgTAAATCCTCATCA	AA	ggggAgCTgggATCCAgCAgggCTg	78	CTgACAgAAACAACCgCCgAgAACT	AA	ATCATCCAgTAAACCgCC

## S4.5 Split-initiator probes for Figures 4 and S12

Organism: G. gallus domesticus

#### Target mRNA: forkhead box D3 (FoxD3)

#### Probe set: 12 split-initiator probe pairs (each probe carries half an HCR initiator)

HCR amplifier: **B4-Alexa488** 

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCAACCTACCTCCAAC	AA	CgCCgCCCgATAgAgTCATCCCCgC	2	gCgCggTCTggCCggACATATCgCT	AT	TCTCACCATATTCgCTTC
3	CCTCAACCTACCTCCAAC	AA	CgTCgATATCCACgTCCTCggCCgC	4	CCggCgCgTCgTCTCCCTCgCCCAC	AT	TCTCACCATATTCgCTTC
5	CCTCAACCTACCTCCAAC	AA	CCCgCAggTgCTCCTgCTggTgCCg	6	AgCCCTgCATCATgAgCgCCgTCTg	AT	TCTCACCATATTCgCTTC
7	CCTCAACCTACCTCCAAC	AA	AgggCCCggCCAgCCCgTAggCgCC	8	CggggggCAgCCCgTAggggCggCC	AT	TCTCACCATATTCgCTTC
9	CCTCAACCTACCTCCAAC	AA	gCAgAgCggCggggTgCgggTAggC	10	gCCCgACgggCgggATgTAggggTA	AT	TCTCACCATATTCgCTTC
11	CCTCAACCTACCTCCAAC	AA	gCAgCgggCACgCgggCggCAgCAT	12	CTTTgCggCTCAgCTCgCCCgACgg	AT	TCTCACCATATTCgCTTC
13	CCTCAACCTACCTCCAAC	AA	ggCTgggCCCgAgCTgCgCgTTgAA	14	CCCCCAAACTgCTgAgCTgCAgCTg	AT	TCTCACCATATTCgCTTC
15	CCTCAACCTACCTCCAAC	AA	gCTCggATTTCACgATggAgCCCgC	16	CgATgCTgAACgAggggCggCTgCT	AT	TCTCACCATATTCgCTTC
17	CCTCAACCTACCTCCAAC	AA	TggCggCggggGCCgCCgATgATgTT	18	ggAAAgTCTgCgCgCTgggCgCCgA	AT	TCTCACCATATTCgCTTC
19	CCTCAACCTACCTCCAAC	AA	CCgACTgCACggTgACgggCggCCg	20	ACgCCAgCggCTggTgCgCCACCAg	AT	TCTCACCATATTCgCTTC
21	CCTCAACCTACCTCCAAC	AA	gCgCgATggCCgCggTggTCCTggC	22	TgATgTTggTAggCACgCTgAggAT	AT	TCTCACCATATTCgCTTC
23	CCTCAACCTACCTCCAAC	AA	CCgTTTCCCAgAgATACgTCCgggg	24	АААТАААААССССдАААдСдАССТС	AT	TCTCACCATATTCgCTTC

#### Organism: *G. gallus domesticus* Target mRNA: **diencephalon/mesencephalon homeobox 1** (*Dmbx1*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B1-Alexa514**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gAggAgggCAgCAAACgg	AA	CgCTCAgTgAgTTCATggCATgCAg	2	CTgCCTgCTggTgCAggTTgTACAT	TA	gAAgAgTCTTCCTTTACg
5	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	6	ggTgCTgggATCCATAACgTgCCTC	TA	gAAgAgTCTTCCTTTACg
9	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	10	gCTTCTgCAgCTgCTCCTTCTggAg	TA	gAAgAgTCTTCCTTTACg
13	gAggAgggCAgCAAACgg	AA	CAgTgTCAggTATTgTggACTgggT	14	CCTCACTTgggATgCTCTgAgTCTT	TA	gAAgAgTCTTCCTTTACg
17	gAggAgggCAgCAAACgg	AA	CCTCCTCTCTgTCAgTCTggTCCTC	18	TAgCCTCATCCAAggTgCTCTTAAA	TA	gAAgAgTCTTCCTTTACg
21	gAggAgggCAgCAAACgg	AA	CgCTgATgggAgACTCTgATTTTgg	22	CACTgCTAgATgAAggAgTCACggT	TA	gAAgAgTCTTCCTTTACg
25	gAggAgggCAgCAAACgg	AA	CTgCCATgTgCTggCggAATTgCTC	26	AggAgTAATggACCAAgTTgTTggT	TA	gAAgAgTCTTCCTTTACg
29	gAggAgggCAgCAAACgg	AA	gACAgTgCAgggAgCTCAgAggCgC	30	CgTgggAgAgAgATTggTAgTACgA	TA	gAAgAgTCTTCCTTTACg
33	gAggAgggCAgCAAACgg	AA	TCTCAATACTTgTTgTTTTACTgTT	34	CATgCTgCTTTgCCCggAgCCTTAg	TA	gAAgAgTCTTCCTTTACg
37	gAggAgggCAgCAAACgg	AA	gACCTCCggTgCATCTTCTTATggg	38	ggTTCCAggAgTgACATgTCTggTg	TA	gAAgAgTCTTCCTTTACg
41	gAggAgggCAgCAAACgg	AA	CggTgCTAgTAAgACATTAgTAAAT	42	AgCCAgTAgCAgTgTCTgATgCAAT	TA	gAAgAgTCTTCCTTTACg
45	gAggAgggCAgCAAACgg	AA	TCACAgCAgTCCAAAgggACAgTTC	46	gCTCTTTCTgAATgTTACAggCTTA	TA	gAAgAgTCTTCCTTTACg
49	gAggAgggCAgCAAACgg	AA	gTTTTCCCAAAgAATgCATCgACAA	50	TATgTACAAgACAAAgCAggACTCT	TA	gAAgAgTCTTCCTTTACg
53	gAggAgggCAgCAAACgg	AA	ggggAATAAAAgCAAAAgAggCCAC	54	gACTAgCTACCAAAACTgAgAgAgA	TA	gAAgAgTCTTCCTTTACg
57	gAggAgggCAgCAAACgg	AA	TTTgCTCTAAgCACCATTAAgACTC	58	gAgCAgTgAATTgCATAATggTTTT	TA	gAAgAgTCTTCCTTTACg
61	gAggAgggCAgCAAACgg	AA	ggAAgTgCTTAAACAggAAATTCAC	62	AgTAAAggAAAAAACACTTgCCTTT	TA	gAAgAgTCTTCCTTTACg
65	gAggAgggCAgCAAACgg	AA	AATTTggCTTTCATTTTTCTCCCCA	66	AACAATCAAgTCAAAAgTAACCATg	TA	gAAgAgTCTTCCTTTACg
69	gAggAgggCAgCAAACgg	AA	CTAgACCAAAATgCTCTCCAAAAAg	70	AgTTTTTATTgTTCTCTATTgTCgA	TA	gAAgAgTCTTCCTTTACg
73	gAggAgggCAgCAAACgg	AA	TAAgAACAgCTTgCATTAATCgTgg	74	TAgAATTTggTgATCggAgCgTTTT	TA	gAAgAgTCTTCCTTTACg
77	gAggAgggCAgCAAACgg	AA	CTTggCCTCCAgCATTgCAgCATTT	78	AATAGAAAgCCCCCgATTATCACCC	TA	gAAgAgTCTTCCTTTACg

#### Organism: *G. gallus domesticus* Target mRNA: **SRY (sex determining region Y)-box 10** (*Sox10*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B3-Alexa546**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	gTCTTCCTCATCTAgAAggCCAATA	2	CCATggACCCgTCACTCCATgTCTT	TT	CCACTCAACTTTAACCCg
3	gTCCCTgCCTCTATATCT	ΤT	TgTgAATCTTAggCAggACTgCTgC	4	TCCAgCAgggATCAAgATTCATgCA	ΤT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	ΤT	gCTgACAgTgCAgTTCCTgAATCCT	6	ACgggCTATgAAATgAgAAAggCTA	ΤT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	TT	TgCTggAggAgCAAggACCTggTCT	8	TTCACgTTTTCAgCAgACACAgTCA	TT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	ΤT	CAAgTgCATggTAgCTTTCTTggTg	10	AATATTggAACCACATCTgggTgTT	ΤT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	ΤT	AgCTgTgAAAATCAgCAAggAAgCA	12	gAggCggggAgAAAAgCTATAgCgT	ΤT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	ΤT	gggATTAAACAgATgggACAggggg	14	TTATAAAATCCATgCAggAAggggT	ΤT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	ΤT	AgCAgTgATgTACACCCCATCggCC	16	AgATggCgATAATgTgATgAACAAA	ΤT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	ΤT	TgAgTgAAAgTAggAgggAggTgCT	18	TATATCCACgAgAgTATCTTTCCAT	ΤT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	CTCTCTgATCAgTTgTCAggAgTCA	20	TACCCgTTAgAAggTCCCACAACAC	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	gAACCCACTgggCTCATCTCCACCT	22	CCCggCgAgAggCAgTggTggTCTT	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	ΤT	TTCCgCCTCCgTggCTggTACTTgT	24	CCCTCgCCCTgCgTggCCTTgCCAT	ΤT	CCACTCAACTTTAACCCg
25	gTCCCTgCCTCTATATCT	ΤT	AgCCCCgCCAgCCTCCCCCTCCACC	26	ATTCTTgTAgTgAgCCTggATAgAg	ΤT	CCACTCAACTTTAACCCg
27	gTCCCTgCCTCTATATCT	ΤT	CCCAggATgCCTgTggTCCAggTgg	28	gTgCCCATCggACATgggTgACCCT	ΤT	CCACTCAACTTTAACCCg
29	gTCCCTgCCTCTATATCT	TT	gCCgTggCTCTgACCTgAAgAgTgC	30	CTTAggggTggTgggAggAgTgggA	TT	CCACTCAACTTTAACCCg
31	gTCCCTgCCTCTATATCT	ΤT	gAgTCAgCTTTgCCTgCCTgCAgCT	32	TCCCCAAgggAACgCCCTTCTCgCT	ΤT	CCACTCAACTTTAACCCg
33	gTCCCTgCCTCTATATCT	ΤT	ggTggCAggTATTggTCAAATTCgT	34	TggCCTgggTggCCggCgTgTCCgT	ΤT	CCACTCAACTTTAACCCg
35	gTCCCTgCCTCTATATCT	ΤT	AgTgTCCACTggCCgCAgCCAgggC	36	CTCCATgCTgCTTggAgATCCAggC	ΤT	CCACTCAACTTTAACCCg
37	gTCCCTgCCTCTATATCT	ΤT	CCCAgTgCCCCCCTgTTCTCCCTCC	38	AAACCCTgTgAAggCTgCAgCTCCT	ΤT	CCACTCAACTTTAACCCg
39	gTCCCTgCCTCTATATCT	ΤT	TggCTgACCgCTTCACggATgCAgA	40	AgggTCCAgTCATAgCCgCTCAgCA	ΤT	CCACTCAACTTTAACCCg

#### Organism: *G. gallus domesticus* Target mRNA: **EPH receptor A4** (*EphA4*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa647**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCgTAAATCCTCATCA	AA	gCgCCgACggggACCCCggCCATgC	2	CAgACgCCgACgAggAgCgggAggA	AA	ATCATCCAgTAAACCgCC
5	CCTCgTAAATCCTCATCA	AA	CCCAgCTCCCCCTgCACCgAgCggg	6	CCTCCTTCCAgCgggCTCgCgATCC	AA	ATCATCCAgTAAACCgCC
9	CCTCgTAAATCCTCATCA	AA	TgACTgggCTCCATCACATTgCAAA	10	ATCCAATCAgTTCgTAACCAATTAT	AA	ATCATCCAgTAAACCgCC
13	CCTCgTAAATCCTCATCA	AA	TCCTTgTCgTTgTTTgATTCATAgT	14	gCAAACTggCTCTCTCgAATAAAAC	AA	ATCATCCAgTAAACCgCC
17	CCTCgTAAATCCTCATCA	AA	AgTgggCACTTCTTATAgAAgACAC	18	ggAAACTgTgCCAggTTTCgAACTg	AA	ATCATCCAgTAAACCgCC
21	CCTCgTAAATCCTCATCA	AA	ACgTCCTTCTCTTCCgAgTTgTTgA	22	CCATCTgCCCCgCAgTACATTTTTg	AA	ATCATCCAgTAAACCgCC
25	CCTCgTAAATCCTCATCA	AA	ATTTTgCAAgCTTggCATTCACCAT	26	TCTgTTgAgAgCgCCTTgTAgTATC	AA	ATCATCCAgTAAACCgCC
29	CCTCgTAAATCCTCATCA	AA	AAgAAgCCCCgATCACAggTgCAgg	30	ATggATgCAgCATCATTTTCTgCTC	AA	ATCATCCAgTAAACCgCC
33	CCTCgTAAATCCTCATCA	AA	gCgCTCCACTCCAAgTTCACTgACg	34	TCgTCCCgTCCTCCCTTgTTCTgTg	AA	ATCATCCAgTAAACCgCC
37	CCTCgTAAATCCTCATCA	AA	AAATgTACACCACTgCCACAggACC	38	gTTTTCAgCCCgTTCTgCTgggggC	AA	ATCATCCAgTAAACCgCC
41	CCTCgTAAATCCTCATCA	AA	TgCTTggACACTCCATTCACTgCCC	42	gACACAgCTTggTCCTggCTggggT	AA	ATCATCCAgTAAACCgCC
45	CCTCgTAAATCCTCATCA	AA	gCAACgCTgTgCCTCgTTATCTCTT	46	ggCCTgTCAggTTCCAgCCAggCCA	AA	ATCATCCAgTAAACCgCC
49	CCTCgTAAATCCTCATCA	AA	gCTgTCTTCACAATgCgATAgCTgC	50	AAACCTTTgATgTCAgTATTCCTgg	AA	ATCATCCAgTAAACCgCC
53	CCTCgTAAATCCTCATCA	AA	AACTCAAACggCCCACTgAAgTCTC	54	ATgggggAAggAACTgTgTTAgTTg	AA	ATCATCCAgTAAACCgCC
57	CCTCgTAAATCCTCATCA	AA	gCTgCAATgAgAATgACCACgAgAA	58	TTgCTgCgCCTCCTgCTgATgACAA	AA	ATCATCCAgTAAACCgCC
61	CCTCgTAAATCCTCATCA	AA	TATGTAAAAggATCCACATATGTTC	62	TCCCTCACAgCTTgATTTggATCCT	AA	ATCATCCAgTAAACCgCC
65	CCTCgTAAATCCTCATCA	AA	CATTTAGTAACAACgCCTTCCAAgT	66	TACTCAgTTATgATCATTACTggTT	AA	ATCATCCAgTAAACCgCC
69	CCTCgTAAATCCTCATCA	AA	TCCCgATgCACATAgCTCATgTCAg	70	TTgACCAgTATgTTTCgAgCAgCTA	AA	ATCATCCAgTAAACCgCC
73	CCTCgTAAATCCTCATCA	AA	CCAAATTTAggTCTgTCgCTgCgTT	74	AgTTTgTCCAgCATgTTgACAATCT	AA	ATCATCCAgTAAACCgCC
77	CCTCgTAAATCCTCATCA	AA	ggggAgCTgggATCCAgCAgggCTg	78	CTgACAgAAACAACCgCCgAgAACT	AA	ATCATCCAgTAAACCgCC

## S4.6 Split-initiator probes for Figures S13–S17

Organism: G. gallus domesticus

#### Target mRNA: **EPH receptor A4** (*EphA4*)

Probe set: 20 split-initiator probe pairs (each probe carries half an HCR initiator)

HCR amplifier: **B1-Alexa546** 

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	gAggAgggCAgCAAACgg	AA	TACACgCgggAgCCggTgACggCCC	4	TCCAgCAgggTCACTTCgTTggCgg	TA	gAAgAgTCTTCCTTTACg
7	gAggAgggCAgCAAACgg	AA	TCATCCATTATgCTCACTTCCTCCC	8	TggTAggTgCggATCggAgTgTTCT	TA	gAAgAgTCTTCCTTTACg
11	gAggAgggCAgCAAACgg	AA	TATACCCTCTgAgCCCCCTCgCggg	12	TCTCTCAgCgTgAACTTgATTTCAA	TA	gAAgAgTCTTCCTTTACg
15	gAggAgggCAgCAAACgg	AA	ACCTCTgTATTCAgCTTCATgATCC	16	TTCTTgCTgAgAggCCCCACgTCCC	TA	gAAgAgTCTTCCTTTACg
19	gAggAgggCAgCAAACgg	AA	gATgTATCAgCCCCAgTAATggTgT	20	CAggAgCCACgAACCTCCACCAgAg	TA	gAAgAgTCTTCCTTTACg
23	gAggAgggCAgCAAACgg	AA	CAgTTgCCAATgggTACCAgCCATT	24	CgTTCTTCATAgCCAgCATTgCACA	TA	gAAgAgTCTTCCTTTACg
27	gAggAgggCAgCAAACgg	AA	TgAggCgggCATTTggCACATgCAA	28	gTAgAgCCTTCCCAgATggAgTAgC	TA	gAAgAgTCTTCCTTTACg
31	gAggAgggCAgCAAACgg	AA	ggTgCggATggAgggCgAgTgCAgg	32	TCgTTgACgTTggAAATCAggTTCT	TA	gAAgAgTCTTCCTTTACg
35	gAggAgggCAgCAAACgg	AA	CgCTTgCACACCACgTTgTAggAgA	36	CAgTggCTgggCTCCCCTgCCCCgC	TA	gAAgAgTCTTCCTTTACg
39	gAggAgggCAgCAAACgg	AA	AggAggTCAgTgATggAAACCTTCg	40	ACCTCAAAggTgTAgTTggTgTgTg	TA	gAAgAgTCTTCCTTTACg
43	gAggAgggCAgCAAACgg	AA	gCTgCTTggTTAgTTgTCACAgTgA	44	gCCTggATCAATgCAATTggggATg	TA	gAAgAgTCTTCCTTTACg
47	gAggAgggCAgCAAACgg	AA	ACTTCATACTCCAggATgACTCCAT	48	TCgTTTTggTCCTTTTCATAgTACT	TA	gAAgAgTCTTCCTTTACg
51	gAggAgggCAgCAAACgg	AA	TgAAATACATATgAAgTCAgggggT	52	TATCCTgCTgCTgTCCTggCCCgCA	TA	gAAgAgTCTTCCTTTACg
55	gAggAgggCAgCAAACgg	AA	ACTgTgggATTggTACCATCgCCAA	56	ACACTgCCAgCCACTgAAACAAgCA	TA	gAAgAgTCTTCCTTTACg
59	gAggAgggCAgCAAACgg	AA	TCTgCCTCTTgCTTAgCTTTACTgT	60	ACACCTTggTTCAAATgTTTCTCCT	TA	gAAgAgTCTTCCTTTACg
63	gAggAgggCAgCAAACgg	AA	TTCAgAgTCTTgATAgCCACACAgA	64	CTCCgTTgTTTgTCAgTgTAACCAg	TA	gAAgAgTCTTCCTTTACg
67	gAggAgggCAgCAAACgg	AA	CgAAgCATCCCCACCAACTggATTA	68	AgATACTTCATTCCTgAgCCgATgC	TA	gAAgAgTCTTCCTTTACg
71	gAggAgggCAgCAAACgg	AA	TggAgAgCAATggggCAgTCCATTg	72	TTCTgCCAgCAgTCTAACATCAgCT	TA	gAAgAgTCTTCCTTTACg
75	gAggAgggCAgCAAACgg	AA	CTCTTCAggCTgTTAgggTTgCggA	76	CTgggTCTggAgCTCTCgCTgCCTg	TA	gAAgAgTCTTCCTTTACg
79	gAggAgggCAgCAAACgg	AA	TCCATTTTAATggCTTggAgCCAgT	80	gCAgCTgTgAAgTTATCCTTgTATC	TA	gAAgAgTCTTCCTTTACg

#### Organism: *G. gallus domesticus* Target mRNA: **early growth response 2** (*Egr2*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B3-Alexa647**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	gCAgTgCTgggCTgCTTgCAgCTgT	2	CgTgCCAgggCTgTgCCAgggCTCA	TT	CCACTCAACTTTAACCCg
3	gTCCCTgCCTCTATATCT	TT	gTCTggCggTAACTATTTATggggg	4	CCCgCCgCAgCTCgCgCTggAggAC	TT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	TT	CACgCCgCTCATCTggTCgAACggC	6	gTCCACgCCgAgCATgCCgTCTgCg	TT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	TT	AACCAACCCCAACCAgTgCgTACAA	8	CTCTTCggTCACCgTAAAgACAAAA	TT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	TT	CCACCCCCCCCCCCGCAgACgCAA	10	gTgCgCTCgTCCAgCCCgggCCCTT	TT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	TT	ggTgTCAgCgTggAATAATTAAgAg	12	CCCCCggCAggCACCTACggAAATA	TT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	TT	TgCACTTgAgTAgCTgAgAgCCTgA	14	ggAggCCgCgAgCAgAgCCTTggCT	TT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	TT	TTTTgCCTggCAgCCCAAATggTgC	16	TgTgACAAgTgTggTAACgCgACTT	TT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	TT	CgACCAgTCATTACTTTCCTCCgCA	18	CAGAATAAATACgggATATCTCACC	TT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	ATATAAggACTgAggAACggggCCC	20	CCCTgAATgCCCgggACgTCACTgC	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	CgCgCCCTTgCgCTCCTTCTggCgC	22	TCCgggCTgCggggCggCggCggTC	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	TT	ACCCCCCACATgCAgCCgggTACgg	24	gCTgTCgTCggAgCggATCggTAAg	TT	CCACTCAACTTTAACCCg
25	gTCCCTgCCTCTATATCT	TT	ggCAAACTTTCCggTCTCggCgTCg	26	ACAgCCATACTACAACCAgggAggg	ΤT	CCACTCAACTTTAACCCg
27	gTCCCTgCCTCTATATCT	TT	CgCTAAgATgAggggAggCgAAAgC	28	TgTgggACCAggTggCAAAgCTgCC	TT	CCACTCAACTTTAACCCg
29	gTCCCTgCCTCTATATCT	TT	CgCTgTCCCTTCgggAgCCTgggAA	30	CCATgTgCCACCTCTCCggCAgACg	TT	CCACTCAACTTTAACCCg
31	gTCCCTgCCTCTATATCT	TT	gAggTTgTgCTCCgCggCCgAgACA	32	TgCCgACTgAggACTCCACgACTCT	ΤT	CCACTCAACTTTAACCCg
33	gTCCCTgCCTCTATATCT	TT	gTCgCggCCAggCCgATgTgCgggg	34	АААААААААААдддадаааадса	ΤT	CCACTCAACTTTAACCCg
35	gTCCCTgCCTCTATATCT	TT	AgCCATggTTATCCAAggCTgTggC	36	TgATgCACgACgCTCCggCTgTgAC	TT	CCACTCAACTTTAACCCg
37	gTCCCTgCCTCTATATCT	TT	CTCACCACgACggCACCgCATgCAT	38	TCACACCACAAggCACCAAggAC	TT	CCACTCAACTTTAACCCg
39	gTCCCTgCCTCTATATCT	TT	gAgCgCCgACCCTgCCgTCgAAggg	40	TTATTCgTCgTCgCTTATAAACgCC	ΤT	CCACTCAACTTTAACCCg

## S4.7 Split-initiator probes Figures 5, S18, and S19

Organism: G. gallus domesticus

Target mRNA: diencephalon/mesencephalon homeobox 1 (*Dmbx1*) Probe set: 20 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B1-Alexa546

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gAggAgggCAgCAAACgg	AA	CgCTCAgTgAgTTCATggCATgCAg	2	CTgCCTgCTggTgCAggTTgTACAT	TA	gAAgAgTCTTCCTTTACg
5	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	6	ggTgCTgggATCCATAACgTgCCTC	TA	gAAgAgTCTTCCTTTACg
9	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	10	gCTTCTgCAgCTgCTCCTTCTggAg	TA	gAAgAgTCTTCCTTTACg
13	gAggAgggCAgCAAACgg	AA	CAgTgTCAggTATTgTggACTgggT	14	CCTCACTTgggATgCTCTgAgTCTT	TA	gAAgAgTCTTCCTTTACg
17	gAggAgggCAgCAAACgg	AA	CCTCCTCTCTgTCAgTCTggTCCTC	18	TAgCCTCATCCAAggTgCTCTTAAA	TA	gAAgAgTCTTCCTTTACg
21	gAggAgggCAgCAAACgg	AA	CgCTgATgggAgACTCTgATTTTgg	22	CACTgCTAgATgAAggAgTCACggT	TA	gAAgAgTCTTCCTTTACg
25	gAggAgggCAgCAAACgg	AA	CTgCCATgTgCTggCggAATTgCTC	26	AggAgTAATggACCAAgTTgTTggT	TA	gAAgAgTCTTCCTTTACg
29	gAggAgggCAgCAAACgg	AA	gACAgTgCAgggAgCTCAgAggCgC	30	CgTgggAgAgAgATTggTAgTACgA	TA	gAAgAgTCTTCCTTTACg
33	gAggAgggCAgCAAACgg	AA	TCTCAATACTTgTTgTTTTACTgTT	34	CATgCTgCTTTgCCCggAgCCTTAg	TA	gAAgAgTCTTCCTTTACg
37	gAggAgggCAgCAAACgg	AA	gACCTCCggTgCATCTTCTTATggg	38	ggTTCCAggAgTgACATgTCTggTg	TA	gAAgAgTCTTCCTTTACg
41	gAggAgggCAgCAAACgg	AA	CggTgCTAgTAAgACATTAgTAAAT	42	AgCCAgTAgCAgTgTCTgATgCAAT	TA	gAAgAgTCTTCCTTTACg
45	gAggAgggCAgCAAACgg	AA	TCACAgCAgTCCAAAgggACAgTTC	46	gCTCTTTCTgAATgTTACAggCTTA	TA	gAAgAgTCTTCCTTTACg
49	gAggAgggCAgCAAACgg	AA	gTTTTCCCAAAgAATgCATCgACAA	50	TATgTACAAgACAAAgCAggACTCT	TA	gAAgAgTCTTCCTTTACg
53	gAggAgggCAgCAAACgg	AA	ggggAATAAAAgCAAAAgAggCCAC	54	gACTAgCTACCAAAACTgAgAgAgA	TA	gAAgAgTCTTCCTTTACg
57	gAggAgggCAgCAAACgg	AA	TTTgCTCTAAgCACCATTAAgACTC	58	gAgCAgTgAATTgCATAATggTTTT	TA	gAAgAgTCTTCCTTTACg
61	gAggAgggCAgCAAACgg	AA	ggAAgTgCTTAAACAggAAATTCAC	62	AgTAAAggAAAAAACACTTgCCTTT	TA	gAAgAgTCTTCCTTTACg
65	gAggAgggCAgCAAACgg	AA	AATTTggCTTTCATTTTTCTCCCCA	66	AACAATCAAgTCAAAAgTAACCATg	TA	gAAgAgTCTTCCTTTACg
69	gAggAgggCAgCAAACgg	AA	CTAgACCAAAATgCTCTCCAAAAAg	70	AgTTTTTATTgTTCTCTATTgTCgA	TA	gAAgAgTCTTCCTTTACg
73	gAggAgggCAgCAAACgg	AA	TAAgAACAgCTTgCATTAATCgTgg	74	TAgAATTTggTgATCggAgCgTTTT	TA	gAAgAgTCTTCCTTTACg
77	gAggAgggCAgCAAACgg	AA	CTTggCCTCCAgCATTgCAgCATTT	78	AATAgAAAgCCCCCgATTATCACCC	TA	gAAgAgTCTTCCTTTACg

#### Organism: *G. gallus domesticus* Target mRNA: **diencephalon/mesencephalon homeobox 1** (*Dmbx1*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa647**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	CCTCgTAAATCCTCATCA	AA	AgTCgggCgCgTgCTgTgCTTgCTg	4	gggTgAgggCgTgCACTgAgggCCg	AA	ATCATCCAgTAAACCgCC
7	CCTCgTAAATCCTCATCA	AA	AggCCgTgCgACTTCgCCTTTgTTT	8	CCAgTgCCTCCAgCTgTTgggCAgT	AA	ATCATCCAgTAAACCgCC
11	CCTCgTAAATCCTCATCA	AA	CACTgTggCTgCTTTCACAgTCCTT	12	CCAgCACAggTggTTCCgTCTTCCC	AA	ATCATCCAgTAAACCgCC
15	CCTCgTAAATCCTCATCA	AA	ACAgggTTAggTTgAggTCTgTgCT	16	gCgCTgATTCACTggCTgACTgCTC	AA	ATCATCCAgTAAACCgCC
19	CCTCgTAAATCCTCATCA	AA	CgTCCACACCAggACTCTTgTCCAC	20	TCgCTCTCTTgCAgTTCAAAgCCTT	AA	ATCATCCAgTAAACCgCC
23	CCTCgTAAATCCTCATCA	AA	AggAgTAggAgTgAgTTTgAgCCAg	24	gCAggCggAAgAggCTCAggggTgA	AA	ATCATCCAgTAAACCgCC
27	CCTCgTAAATCCTCATCA	AA	TggCAgACggTgTCCCCATTTCgAA	28	TgTTgACATTCATgCCCAAgTAggg	AA	ATCATCCAgTAAACCgCC
31	CCTCgTAAATCCTCATCA	AA	ggATgggTgAggACCAgACCTgCTg	32	ggCTTggAAgAgAgCTggAggCCTg	AA	ATCATCCAgTAAACCgCC
35	CCTCgTAAATCCTCATCA	AA	gTAgggTgTCAAgCCCAAgggATgC	36	ACggTTgCCCATCTggCAgTCAgTT	AA	ATCATCCAgTAAACCgCC
39	CCTCgTAAATCCTCATCA	AA	ACTCCAggAAgAgATgAgggTggAA	40	AAAgTTTTCCCTgATAgggAgCACC	AA	ATCATCCAgTAAACCgCC
43	CCTCgTAAATCCTCATCA	AA	TCCTCCTCAAATATTTAAAgAAgAC	44	CTgTCTAAACACACATCCTCTCCCT	AA	ATCATCCAgTAAACCgCC
47	CCTCgTAAATCCTCATCA	AA	ACATTATCgCAgggATgAggTgAgg	48	AAAAAgggTgTATATAACACggTTg	AA	ATCATCCAgTAAACCgCC
51	CCTCgTAAATCCTCATCA	AA	gCggTggATgCTTTCAACATTgTAA	52	TTCTgTAACACTgACAgTAACACAC	AA	ATCATCCAgTAAACCgCC
55	CCTCgTAAATCCTCATCA	AA	gggAgCgTggCTgATTTgTgACTTT	56	ААСССААдААдАдСААСТАдСТдТд	AA	ATCATCCAgTAAACCgCC
59	CCTCgTAAATCCTCATCA	AA	TCAgCTTTAgCAgAgAAgAgAgAAg	60	TTgCATCATTTCCTggCCgTTATAA	AA	ATCATCCAgTAAACCgCC
63	CCTCgTAAATCCTCATCA	AA	TCTgTCTgTgAACAAgTgCTATTAg	64	CAgCAgCATTTggCCAgCATTTTgT	AA	ATCATCCAgTAAACCgCC
67	CCTCgTAAATCCTCATCA	AA	TTACACTTCACTgAAgACCAAAgAg	68	AACCCCATAATTTgTAAATgggggA	AA	ATCATCCAgTAAACCgCC
71	CCTCgTAAATCCTCATCA	AA	gTATgAACACAgTgggAgTTCATAC	72	TTgTCAAggAACCATATAATTCATg	AA	ATCATCCAgTAAACCgCC
75	CCTCgTAAATCCTCATCA	AA	AgggTCTgAAgCTgCACAgCTTgAg	76	CACTTGTTACATTCTCACTTGCTAA	AA	ATCATCCAgTAAACCgCC
79	CCTCgTAAATCCTCATCA	AA	CAAgCCAATCTACTCCTCgCTgCAg	80	ggTTgCTTggggACATggTACTTTT	AA	ATCATCCAgTAAACCgCC

#### Organism: *G. gallus domesticus* Target mRNA: **EPH receptor A4** (*EphA4*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B1-Alexa546**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	gAggAgggCAgCAAACgg	AA	TACACgCgggAgCCggTgACggCCC	4	TCCAgCAgggTCACTTCgTTggCgg	TA	gAAgAgTCTTCCTTTACg
7	gAggAgggCAgCAAACgg	AA	TCATCCATTATgCTCACTTCCTCCC	8	TggTAggTgCggATCggAgTgTTCT	TA	gAAgAgTCTTCCTTTACg
11	gAggAgggCAgCAAACgg	AA	TATACCCTCTgAgCCCCCTCgCggg	12	TCTCTCAgCgTgAACTTgATTTCAA	TA	gAAgAgTCTTCCTTTACg
15	gAggAgggCAgCAAACgg	AA	ACCTCTgTATTCAgCTTCATgATCC	16	TTCTTgCTgAgAggCCCCACgTCCC	TA	gAAgAgTCTTCCTTTACg
19	gAggAgggCAgCAAACgg	AA	gATgTATCAgCCCCAgTAATggTgT	20	CAggAgCCACgAACCTCCACCAgAg	TA	gAAgAgTCTTCCTTTACg
23	gAggAgggCAgCAAACgg	AA	CAgTTgCCAATgggTACCAgCCATT	24	CgTTCTTCATAgCCAgCATTgCACA	TA	gAAgAgTCTTCCTTTACg
27	gAggAgggCAgCAAACgg	AA	TgAggCgggCATTTggCACATgCAA	28	gTAgAgCCTTCCCAgATggAgTAgC	TA	gAAgAgTCTTCCTTTACg
31	gAggAgggCAgCAAACgg	AA	ggTgCggATggAgggCgAgTgCAgg	32	TCgTTgACgTTggAAATCAggTTCT	TA	gAAgAgTCTTCCTTTACg
35	gAggAgggCAgCAAACgg	AA	CgCTTgCACACCACgTTgTAggAgA	36	CAgTggCTgggCTCCCCTgCCCCgC	TA	gAAgAgTCTTCCTTTACg
39	gAggAgggCAgCAAACgg	AA	AggAggTCAgTgATggAAACCTTCg	40	ACCTCAAAggTgTAgTTggTgTgTg	TA	gAAgAgTCTTCCTTTACg
43	gAggAgggCAgCAAACgg	AA	gCTgCTTggTTAgTTgTCACAgTgA	44	gCCTggATCAATgCAATTggggATg	TA	gAAgAgTCTTCCTTTACg
47	gAggAgggCAgCAAACgg	AA	ACTTCATACTCCAggATgACTCCAT	48	TCgTTTTggTCCTTTTCATAgTACT	TA	gAAgAgTCTTCCTTTACg
51	gAggAgggCAgCAAACgg	AA	TgAAATACATATgAAgTCAgggggT	52	TATCCTgCTgCTgTCCTggCCCgCA	TA	gAAgAgTCTTCCTTTACg
55	gAggAgggCAgCAAACgg	AA	ACTgTgggATTggTACCATCgCCAA	56	ACACTgCCAgCCACTgAAACAAgCA	TA	gAAgAgTCTTCCTTTACg
59	gAggAgggCAgCAAACgg	AA	TCTgCCTCTTgCTTAgCTTTACTgT	60	ACACCTTggTTCAAATgTTTCTCCT	TA	gAAgAgTCTTCCTTTACg
63	gAggAgggCAgCAAACgg	AA	TTCAgAgTCTTgATAgCCACACAgA	64	CTCCgTTgTTTgTCAgTgTAACCAg	TA	gAAgAgTCTTCCTTTACg
67	gAggAgggCAgCAAACgg	AA	CgAAgCATCCCCACCAACTggATTA	68	AgATACTTCATTCCTgAgCCgATgC	TA	gAAgAgTCTTCCTTTACg
71	gAggAgggCAgCAAACgg	AA	TggAgAgCAATggggCAgTCCATTg	72	TTCTgCCAgCAgTCTAACATCAgCT	TA	gAAgAgTCTTCCTTTACg
75	gAggAgggCAgCAAACgg	AA	CTCTTCAggCTgTTAgggTTgCggA	76	CTgggTCTggAgCTCTCgCTgCCTg	TA	gAAgAgTCTTCCTTTACg
79	gAggAgggCAgCAAACgg	AA	TCCATTTTAATggCTTggAgCCAgT	80	gCAgCTgTgAAgTTATCCTTgTATC	TA	gAAgAgTCTTCCTTTACg

#### Organism: *G. gallus domesticus* Target mRNA: **EPH receptor A4** (*EphA4*) Probe set: **20 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa647**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCgTAAATCCTCATCA	AA	gCgCCgACggggACCCCggCCATgC	2	CAgACgCCgACgAggAgCgggAggA	AA	ATCATCCAgTAAACCgCC
5	CCTCgTAAATCCTCATCA	AA	CCCAgCTCCCCCTgCACCgAgCggg	6	CCTCCTTCCAgCgggCTCgCgATCC	AA	ATCATCCAgTAAACCgCC
9	CCTCgTAAATCCTCATCA	AA	TgACTgggCTCCATCACATTgCAAA	10	ATCCAATCAGTTCGTAACCAATTAT	AA	ATCATCCAgTAAACCgCC
13	CCTCgTAAATCCTCATCA	AA	TCCTTgTCgTTgTTTgATTCATAgT	14	gCAAACTggCTCTCTCgAATAAAAC	AA	ATCATCCAgTAAACCgCC
17	CCTCgTAAATCCTCATCA	AA	AgTgggCACTTCTTATAgAAgACAC	18	ggAAACTgTgCCAggTTTCgAACTg	AA	ATCATCCAgTAAACCgCC
21	CCTCgTAAATCCTCATCA	AA	ACgTCCTTCTCTTCCgAgTTgTTgA	22	CCATCTgCCCCgCAgTACATTTTTg	AA	ATCATCCAgTAAACCgCC
25	CCTCgTAAATCCTCATCA	AA	ATTTTgCAAgCTTggCATTCACCAT	26	TCTgTTgAgAgCgCCTTgTAgTATC	AA	ATCATCCAgTAAACCgCC
29	CCTCgTAAATCCTCATCA	AA	AAgAAgCCCCgATCACAggTgCAgg	30	ATggATgCAgCATCATTTTCTgCTC	AA	ATCATCCAgTAAACCgCC
33	CCTCgTAAATCCTCATCA	AA	gCgCTCCACTCCAAgTTCACTgACg	34	TCgTCCCgTCCTCCCTTgTTCTgTg	AA	ATCATCCAgTAAACCgCC
37	CCTCgTAAATCCTCATCA	AA	AAATgTACACCACTgCCACAggACC	38	gTTTTCAgCCCgTTCTgCTgggggC	AA	ATCATCCAgTAAACCgCC
41	CCTCgTAAATCCTCATCA	AA	TgCTTggACACTCCATTCACTgCCC	42	gACACAgCTTggTCCTggCTggggT	AA	ATCATCCAgTAAACCgCC
45	CCTCgTAAATCCTCATCA	AA	gCAACgCTgTgCCTCgTTATCTCTT	46	ggCCTgTCAggTTCCAgCCAggCCA	AA	ATCATCCAgTAAACCgCC
49	CCTCgTAAATCCTCATCA	AA	gCTgTCTTCACAATgCgATAgCTgC	50	AAACCTTTgATgTCAgTATTCCTgg	AA	ATCATCCAgTAAACCgCC
53	CCTCgTAAATCCTCATCA	AA	AACTCAAACggCCCACTgAAgTCTC	54	ATgggggAAggAACTgTgTTAgTTg	AA	ATCATCCAgTAAACCgCC
57	CCTCgTAAATCCTCATCA	AA	gCTgCAATgAgAATgACCACgAgAA	58	TTgCTgCgCCTCCTgCTgATgACAA	AA	ATCATCCAgTAAACCgCC
61	CCTCgTAAATCCTCATCA	AA	TATGTAAAAggATCCACATATGTTC	62	TCCCTCACAgCTTgATTTggATCCT	AA	ATCATCCAgTAAACCgCC
65	CCTCgTAAATCCTCATCA	AA	CATTTAGTAACAACgCCTTCCAAgT	66	TACTCAgTTATgATCATTACTggTT	AA	ATCATCCAgTAAACCgCC
69	CCTCgTAAATCCTCATCA	AA	TCCCgATgCACATAgCTCATgTCAg	70	TTgACCAgTATgTTTCgAgCAgCTA	AA	ATCATCCAgTAAACCgCC
73	CCTCgTAAATCCTCATCA	AA	CCAAATTTAggTCTgTCgCTgCgTT	74	AgTTTgTCCAgCATgTTgACAATCT	AA	ATCATCCAgTAAACCgCC
77	CCTCgTAAATCCTCATCA	AA	ggggAgCTgggATCCAgCAgggCTg	78	CTgACAgAAACAACCgCCgAgAACT	AA	ATCATCCAgTAAACCgCC

## S4.8 Split-initiator probes for Figure 6, S20–S28

#### Organism: H. sapiens sapiens

#### Target mRNA: destabilized enhanced green fluorescent protein (Tg(d2eGFP))Probe set: 12 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B3-Alexa594 (Figures 6A and S20)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	TTgTggCCgTTTACgTCgCCgTCCA	2	CCCTCgCCCTCgCCggACACgCTgA	TT	CCACTCAACTTTAACCCg
3	gTCCCTgCCTCTATATCT	TT	AgggTCAgCTTgCCgTAggTggCAT	4	AgCTTgCCggTggTgCAgATgAACT	TT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	TT	gTCACgAgggTgggCCAgggCACgg	6	AAgCACTgCACgCCgTAggTCAggg	TT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	TT	TgCTTCATgTggTCggggTAgCggC	8	ggCATggCggACTTgAAgAAgTCgT	TT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	TT	ATggTgCgCTCCTggACgTAgCCTT	10	TTgTAgTTgCCgTCgTCCTTgAAgA	TT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	TT	CCCTCgAACTTCACCTCggCgCggg	12	AgCTCgATgCggTTCACCAgggTgT	TT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	TT	CCgTCCTCCTTgAAgTCgATgCCCT	14	TACTCCAgCTTgTgCCCCAggATgT	TT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	TT	ATATAgACgTTgTggCTgTTgTAgT	16	ATgCCgTTCTTCTgCTTgTCggCCA	TT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	TT	TTgTggCggATCTTgAAgTTCACCT	18	gCgAgCTgCACgCTgCCgTCCTCgA	TT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	ATgggggTgTTCTgCTggTAgTggT	20	TCgggCAgCAgCACgggggCCgTCgC	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	gCggACTgggTgCTCAggTAgTggT	22	CgCTTCTCgTTgggggTCTTTgCTCA	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	TT	ACgAACTCCAgCAggACCATgTgAT	24	ATgCCgAgAgTgATCCCggCggCgg	TT	CCACTCAACTTTAACCCg

#### Organism: H. sapiens sapiens

#### Target mRNA: glyceraldehyde-3-phosphate dehydrogenase (*GAPDH*) Probe set: 10 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: **B5-Alexa488** (Figures 6B, S24, and S23)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	CTCACTCCCAATCTCTAT	AA	gggTCATTgATggCAACAATATCCA	4	TAAACCATgTAgTTgAggTCAATgA	AA	CTACCCTACAAATCCAAT
7	CTCACTCCCAATCTCTAT	AA	TTTCCATTgATgACAAgCTTCCCgT	8	TCTCgCTCCTggAAgATggTgATgg	AA	CTACCCTACAAATCCAAT
11	CTCACTCCCAATCTCTAT	AA	gCCTTCTCCATggTggTgAAgACgC	12	TTggCTCCCCCTgCAAATgAgCCC	AA	CTACCCTACAAATCCAAT
15	CTCACTCCCAATCTCTAT	AA	AggCTgTTgTCATACTTCTCATggT	16	gTgCAggAggCATTgCTgATgATCT	AA	CTACCCTACAAATCCAAT
19	CTCACTCCCAATCTCTAT	AA	gCATggACTgTggTCATgAgTCCTT	20	TCCACAgTCTTCTgggTggCAgTgA	AA	CTACCCTACAAATCCAAT
23	CTCACTCCCAATCTCTAT	AA	gCCTTggCAgCgCCAgTAgAggCAg	24	TTCAgCTCAgggATgACCTTgCCCA	AA	CTACCCTACAAATCCAAT
27	CTCACTCCCAATCTCTAT	AA	ggTTTTTCTAgACggCAggTCAggT	28	ACCTTCTTgATgTCATCATATTTgg	AA	CTACCCTACAAATCCAAT
31	CTCACTCCCAATCTCTAT	AA	CTgTTgAAgTCAgAggAgACCACCT	32	gCgTCAAAggTggAggAgTgggTgT	AA	CTACCCTACAAATCCAAT
35	CTCACTCCCAATCTCTAT	AA	CTggTggTCCAggggTCTTACTCCT	36	TCTCTTCCTCTTgTgCTCTTgCTgg	AA	CTACCCTACAAATCCAAT
39	CTCACTCCCAATCTCTAT	AA	CTACATggCAACTgTgAggAggggA	40	CCTAggCCCCTCCCCTCTTCAAggg	AA	CTACCCTACAAATCCAAT

#### Organism: H. sapiens sapiens

100

#### Target mRNA: glyceraldehyde-3-phosphate dehydrogenase (*GAPDH*) Probe set: 10 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B4-Alexa594 (Figures 6B and S24), B4-Alexa488 (Figures 6C and S27)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCAACCTACCTCCAAC	AA	ACCAggCgCCCAATACgACCAAATC	2	TTACCAgAgTTAAAAgCAgCCCTgg	AT	TCTCACCATATTCgCTTC
5	CCTCAACCTACCTCCAAC	AA	CCATgggTggAATCATATTggAACA	6	TCAgCCTTgACggTgCCATggAATT	AT	TCTCACCATATTCgCTTC
9	CCTCAACCTACCTCCAAC	AA	gCATCgCCCCACTTgATTTTggAgg	10	gTggACTCCACgACgTACTCAgCgC	AT	TCTCACCATATTCgCTTC
13	CCTCAACCTACCTCCAAC	AA	gCAgAgggggCAgAgATgATgACCC	14	ACACCCATgACgAACATgggggCAT	AT	TCTCACCATATTCgCTTC
17	CCTCAACCTACCTCCAAC	AA	TTggCCAggggTgCTAAgCAgTTgg	18	ACgATACCAAAgTTgTCATggATgA	AT	TCTCACCATATTCgCTTC
21	CCTCAACCTACCTCCAAC	AA	TCACgCCACAgTTTCCCggAggggC	22	ATgATgTTCTggAgAgCCCCgCggC	AT	TCTCACCATATTCgCTTC
25	CCTCAACCTACCTCCAAC	AA	CggAAggCCATgCCAgTgAgCTTCC	26	ACCACTgACACgTTggCAgTggggA	AT	TCTCACCATATTCgCTTC
29	CCTCAACCTACCTCCAAC	AA	AgggggCCCTCCgACgCCTgCTTCA	30	TgCTCAgTgTAgCCCAggATgCCCT	AT	TCTCACCATATTCgCTTC
33	CCTCAACCTACCTCCAAC	AA	TggTCgTTgAgggCAATgCCAgCCC	34	TCATACCAggAAATgAgCTTgACAA	AT	TCTCACCATATTCgCTTC
37	CCTCAACCTACCTCCAAC	AA	CAgggACTCCCCAgCAgTgAgggTC	38	TTCAgTgTggTgggggACTgAgTgT	AT	TCTCACCATATTCgCTTC

#### Organism: *H. sapiens sapiens* Target mRNA: actin beta (*ACTB*) Probe set: 10 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B2-Alexa594 (Figure S23)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCgTAAATCCTCATCA	AA	gCggggCggACgCggTCTCggCggT	2	gATCggCAAAggCgAggCTCTgTgC	AA	ATCATCCAgTAAACCgCC
5	CCTCgTAAATCCTCATCA	AA	gCTggCggCgggTgTggACgggCgg	6	gCgCggCgATATCATCATCCATggT	AA	ATCATCCAgTAAACCgCC
9	CCTCgTAAATCCTCATCA	AA	TgCgCAAgTTAggTTTTgTCAAgAA	10	AAgCCATgCCAATCTCATCTTgTTT	AA	ATCATCCAgTAAACCgCC
13	CCTCgTAAATCCTCATCA	AA	АССААААСААААСАААААААААААА	14	СТдАдТСААдССААААААААААААА	AA	ATCATCCAgTAAACCgCC
17	CCTCgTAAATCCTCATCA	AA	CACCTTCACCgTTCCAgTTTTTAAA	18	gggATgCTCgCTCCAACCgACTgCT	AA	ATCATCCAgTAAACCgCC
21	CCTCgTAAATCCTCATCA	AA	AgTCCTCggCCACATTgTgAACTTT	22	АТТАААААААСААСААТдТдСААТС	AA	ATCATCCAgTAAACCgCC
25	CCTCgTAAATCCTCATCA	AA	ACAACgCATCTCATATTTggAATgA	26	TTTTAggATggCAAgggACTTCCTg	AA	ATCATCCAgTAAACCgCC
29	CCTCgTAAATCCTCATCA	AA	ATTCTCCTTAgAgAgAAgTggggTg	30	TgTgTggACTTgggAgAggACTggg	AA	ATCATCCAgTAAACCgCC
33	CCTCgTAAATCCTCATCA	AA	ACACgAAAgCAATgCTATCACCTCC	34	ТТАААААААТТТТдСАТТАСАТААТ	AA	ATCATCCAgTAAACCgCC
37	CCTCgTAAATCCTCATCA	AA	СААААТАААААдТАТТААддСдАА	38	CACgAAggCTCATCATTCAAAATAA	AA	ATCATCCAgTAAACCgCC

## Organism: *H. sapiens sapiens*

#### Target mRNA: **phosphoglycerate kinase 1** (*PGK1*) Probe set: **18 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B1-Alexa488 (Figures 6B and S25**)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gAggAgggCAgCAAACgg	AA	CCgCCCCTTCCCggCCgCTgCTCTC	2	CTACCgCCCCACACCCCgCCTCCCg	TA	gAAgAgTCTTCCTTTACg
5	gAggAgggCAgCAAACgg	AA	CAACgAgggAgCCgACTgCCgACgT	6	gCTggggAgAgAggTCggTgATTCg	TA	gAAgAgTCTTCCTTTACg
9	gAggAgggCAgCAAACgg	AA	gACTCTCATAACgACCCgCTTCCCT	10	gTTgTTCTTCATAggAACATTgAAg	TA	gAAgAgTCTTCCTTTACg
13	gAggAgggCAgCAAACgg	AA	CTTCCCTTCTTCCTCCACATgAAAg	14	AACCTTgTTCCCAgAAgCATCTTTT	TA	gAAgAgTCTTCCTTTACg
17	gAggAgggCAgCAAACgg	AA	gCCAAAAgCATCATTgACATAgACA	18	CATggAgCTgTgggCTCTgTgAgCA	TA	gAAgAgTCTTCCTTTACg
21	gAggAgggCAgCAAACgg	AA	gCTCTCCAAggCCTTTgCAAAgTAg	22	CAggATggCCAggAAgggTCgCTCT	TA	gAAgAgTCTTCCTTTACg
25	gAggAgggCAgCAAACgg	AA	CTTCTCAgCTTTggACATTAggTCT	26	AACAggCAAggTAATCTTCACACCA	TA	gAAgAgTCTTCCTTTACg
29	gAggAgggCAgCAAACgg	AA	CCAgCCAgCAggTATgCCAgAAgCC	30	gCTTTCAggACCACAgTCCAAgCCC	TA	gAAgAgTCTTCCTTTACg
33	gAggAgggCAgCAAACgg	AA	AgCTTCCCATTCAAATACCCCCACA	34	CATgAgAgCTTTggTTCCCCgggCA	TA	gAAgAgTCTTCCTTTACg
37	gAggAgggCAgCAAACgg	AA	gTACTAAATATTgCTgAgAgCATCC	38	ТдТдСАСАддААСТААААддСАддА	TA	gAAgAgTCTTCCTTTACg
41	gAggAgggCAgCAAACgg	AA	ggCCACTAgCTgAATCTTgACATgg	42	TTAAgggTTCCTggCACTgCATCTC	TA	gAAgAgTCTTCCTTTACg
45	gAggAgggCAgCAAACgg	AA	СТААААААТТСАААТдддАТСТТДА	46	ACTCTAgAATgCACAATggTTTAgT	TA	gAAgAgTCTTCCTTTACg
49	gAggAgggCAgCAAACgg	AA	ТААТСАТААТААССТАСАТСААААд	50	TgCTgAgTAgTgAAACAgTgACAAA	TA	gAAgAgTCTTCCTTTACg
53	gAggAgggCAgCAAACgg	AA	TCAATggACACTTTTATTgTTTACT	54	дАСАддААААААААААААТСАСдд	TA	gAAgAgTCTTCCTTTACg
57	gAggAgggCAgCAAACgg	AA	CTgCCCCACTTCTTgCATTCAgCAA	58	TCTAATTgTCCCATCTCTCCACTgC	TA	gAAgAgTCTTCCTTTACg
61	gAggAgggCAgCAAACgg	AA	СТдАТАААААТААААдТТАдААТАА	62	gACTTTTTAAATTATgATCATgTgT	TA	gAAgAgTCTTCCTTTACg
65	gAggAgggCAgCAAACgg	AA	CAAgAgTTgAAAgTggTCACCTCTg	66	AACATggAggTATATACCTgAAAAA	TA	gAAgAgTCTTCCTTTACg
69	gAggAgggCAgCAAACgg	AA	gAgCCTTCCTCCATggTATgAAATA	70	TgAAgAAgTggAAATATATgTggAA	TA	gAAgAgTCTTCCTTTACg

#### Organism: *H. sapiens sapiens*

#### Target mRNA: **phosphoglycerate kinase 1** (*PGK1*) Probe set: **18 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa594 (Figures 6B, 6C, S25, and S27**)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	CCTCgTAAATCCTCATCA	AA	ACCgCgCgggCAggAACAgggCCCA	4	gCTCCggAggCTTgCAgAATgCggA	AA	ATCATCCAgTAAACCgCC
7	CCTCgTAAATCCTCATCA	AA	gTTAgAAAgCgACATTTTggAAATA	8	AACgTCCAgCTTgTCCAgCgTCAgC	AA	ATCATCCAgTAAACCgCC
11	CCTCgTAAATCCTCATCA	AA	CTTAATCCTCTggTTgTTTgTTATC	12	gCAgAATTTgATgCTTgggACAgCA	AA	ATCATCCAgTAAACCgCC
15	CCTCgTAAATCCTCATCA	AA	ggAgTACTTgTCAggCATgggCACA	16	TTTgAgTTCTACAgCAACTggCTCT	AA	ATCATCCAgTAAACCgCC
19	CCTCgTAAATCCTCATCA	AA	AgCCTTCTgTggCAgATTgACTCCT	20	CAgCTCCTTCTTCATCAAAAACCCA	AA	ATCATCCAgTAAACCgCC
23	CCTCgTAAATCCTCATCA	AA	ATCAAACTTgTCAgCAgTgACAAAg	24	AgTggCTTggCCAgTCTTggCATTC	AA	ATCATCCAgTAAACCgCC
27	CCTCgTAAATCCTCATCA	AA	AgTgACAgCCTCAgCATACTTCTTg	28	ACCATTCCACACAATCTgCTTAgCC	AA	ATCATCCAgTAAACCgCC
31	CCTCgTAAATCCTCATCA	AA	CCTAgAAgTggCTTTCACCACCTCA	32	TCCACCACCTATgATggTgATgCAg	AA	ATCATCCAgTAAACCgCC
35	CCTCgTAAATCCTCATCA	AA	CTCCAAACTggCACCACCCCAgTg	36	CCCAggAAggACTTTACCTTCCAgg	AA	ATCATCCAgTAAACCgCC
39	CCTCgTAAATCCTCATCA	AA	CAgAAAATgCTAAgTTgACTTAggg	40	ggTTTTAgCTAATgCCAAgTggAgA	AA	ATCATCCAgTAAACCgCC
43	CCTCgTAAATCCTCATCA	AA	AgATgAgCTgAgATgCTgTgCAACT	44	AATgTATgCAAATCCAgggTgCAgT	AA	ATCATCCAgTAAACCgCC
47	CCTCgTAAATCCTCATCA	AA	ТТТААСАддСААААТАТАААТАТАТ	48	AACTAAgCTAACACTgCTCACTTTC	AA	ATCATCCAgTAAACCgCC
51	CCTCgTAAATCCTCATCA	AA	TACAAATggAATTTCATCTTgTTTC	52	ATggATCATCAATTTTgTCTCACTA	AA	ATCATCCAgTAAACCgCC
55	CCTCgTAAATCCTCATCA	AA	CTATTCTCACCCTTCCTAACAAAgT	56	TAGACATCTGATCCGTTCCTCAAGA	AA	ATCATCCAgTAAACCgCC
59	CCTCgTAAATCCTCATCA	AA	AggCCCTTgATAAAgAATggACATT	60	gCACTAgCACAATgTCTgCCATAAA	AA	ATCATCCAgTAAACCgCC
63	CCTCgTAAATCCTCATCA	AA	ggCTggggCTTTTTTgTTATAAgCC	64	gAgTgggAATCTTgAATgggAggAA	AA	ATCATCCAgTAAACCgCC
67	CCTCgTAAATCCTCATCA	AA	gTgAACAATATAAgCATATTACTTA	68	ТТТСТТТААААААТААААААААА	AA	ATCATCCAgTAAACCgCC
71	CCTCgTAAATCCTCATCA	AA	AATTgTgACAAAACTATACCgAgAg	72	gTTATgTAgACTTTTgATCTAATCT	AA	ATCATCCAgTAAACCgCC

#### Organism: E. coli

### Target mRNA: enhanced green fluorescent protein (*Tg*(*EGFP*)) Probe set: 12 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B3-Alexa594 (Figures 6A and S22)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	TgAAAAgTTCTTCTCCTTTACgCAT	2	ATTCAACAAgAATTgggACAACTCC	TT	CCACTCAACTTTAACCCg
3	gTCCCTgCCTCTATATCT	TT	ATTTgTgCCCATTAACATCACCATC	4	CACCTTCACCCTCTCCACTgACAgA	TT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	TT	TAAgggTAAgTTTTCCgTATgTTgC	6	gTAgTTTTCCAgTAgTgCAAATAAA	TT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	TT	TAgTgACAAgTgTTggCCATggAAC	8	CAAAgCATTgAACACCATAACCgAA	TT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	TT	gCTgTTTCATATgATCTgggTATCT	10	CgggCATggCACTCTTgAAAAAgTC	TT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	TT	ATATAGTTCTTTCCTGTACATAACC	12	TCTTgTAgTTCCCgTCATCTTTgAA	TT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	TT	CACCTTCAAACTTgACTTCAgCACg	14	TTAACTCgATTCTATTAACAAgggT	TT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	TT	TTCCATCTTCTTTAAAATCAATACC	16	TgTATTCCAATTTgTgTCCAAgAAT	TT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	TT	TgATgTATACATTgTgTgAgTTATA	18	TgATTCCATTCTTTTgTTTgTCTgC	TT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	TgTTgTgTCTAATTTTgAAgTTAAC	20	CTgCTAgTTgAACgCTTCCATCTTC	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	CAATTggAgTATTTTgTTgATAATg	22	TgTCTggTAAAAggACAgggCCATC	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	TT	gggCAgATTgTgTggACAggTAATg	24	CTCTCTTTTCgTTgggATCTTTCgA	TT	CCACTCAACTTTAACCCg

#### Organism: E. coli

#### Target mRNA: **GTP-binding protein chain elongation factor EF-G** (*fusA*) Probe set: **18 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa594 (Figures 6B and S26**)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	CCTCgTAAATCCTCATCA	AA	gTgCgATgggTgTTgTACgAgCCAT	2	gCgCACTgATACCgATgTTACggTA	AA	ATCATCCAgTAAACCgCC
5	CCTCgTAAATCCTCATCA	AA	CATGAACTTCACCGATTTTATggTT	6	CCATCCAgTCCATggTTgCAgCgCC	AA	ATCATCCAgTAAACCgCC
9	CCTCgTAAATCCTCATCA	AA	gCTCATACTgCTTAgCCATACCAgA	10	gggTgTCgATgATgTTgATgCgATg	AA	ATCATCCAgTAAACCgCC
13	CCTCgTAAATCCTCATCA	AA	CAACTgCgCAgTAAACCATTACCgC	14	CggTTTCAgACTgCggCTgAACACC	AA	ATCATCCAgTAAACCgCC
17	CCTCgTAAATCCTCATCA	AA	TCAggAAgTTCgCACCCATgCggTC	18	gACgggTTTTgATCTggTTAACAAC	AA	ATCATCCAgTAAACCgCC
21	CCTCgTAAATCCTCATCA	AA	TCATTTTCACCAggTCAACAACACC	22	ggTCAgCgTCgTTCCAgTTgATAgC	AA	ATCATCCAgTAAACCgCC
25	CCTCgTAAATCCTCATCA	AA	ATTCgATCAggTTCTggTgCCATTC	26	TCAgCTCTTCAgAAgCTTCAgCTgC	AA	ATCATCCAgTAAACCgCC
29	CCTCgTAAATCCTCATCA	AA	TTTCgTTgTTCAgAACgCgCTgACg	30	ACgCAgAACCACAggTTACCAggAT	AA	ATCATCCAgTAAACCgCC
. 33	CCTCgTAAATCCTCATCA	AA	CgTTgATCgCAggTACgTCAACCgg	34	gAgTgTCTTTACCgTCgTCCAggAT	AA	ATCATCCAgTAAACCgCC
37	CCTCgTAAATCCTCATCA	AA	ggTTACCAACAAACgggTCggTAgC	38	CACCggAgTAAACACggAAgAAggT	AA	ATCATCCAgTAAACCgCC
41	CCTCgTAAATCCTCATCA	AA	CgATAgCAgCAgCgATgTCgCCCgC	42	TgTCACCAgTggTTACgTCTTTCAg	AA	ATCATCCAgTAAACCgCC
45	CCTCgTAAATCCTCATCA	AA	TCggTTCAACTgCgATggAgATTAC	46	CCATTTTTTCCTggTCAgCTTTggT	AA	ATCATCCAgTAAACCgCC
49	CCTCgTAAATCCTCATCA	AA	ATTCACgCTTCATACggTCAACgAT	50	gTTTACCTACgTTCgCTTCAACgTT	AA	ATCATCCAgTAAACCgCC
53	CCTCgTAAATCCTCATCA	AA	CACgACCACCAgACTgTTTCgCgTg	54	TgTCgATAACAACATgACCATACTg	AA	ATCATCCAgTAAACCgCC
57	CCTCgTAAATCCTCATCA	AA	ATTCgCCAgggATTACACCACCTTT	58	ggATACCTTTATCAACggCCgggAT	AA	ATCATCCAgTAAACCgCC
61	CCTCgTAAATCCTCATCA	AA	CATggTAAgAACCgAAgTgCAgACg	62	TAAACgCCAgTTCAgAggAgTCAAC	AA	ATCATCCAgTAAACCgCC
65	CCTCgTAAATCCTCATCA	AA	CTTCAACCTTCATgATCggCTCAAg	66	CACCggTgTTCTCTTCCggAgTTTC	AA	ATCATCCAgTAAACCgCC
69	CCTCgTAAATCCTCATCA	AA	CAgCgTggATCTTAACgCCAgTAAC	70	ATCCgAACATTTCAgACAgCggTAC	AA	ATCATCCAgTAAACCgCC

#### Organism: E.coli

#### Target mRNA: **GTP-binding protein chain elongation factor EF-G** *fusA* Probe set: **18 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B3-Alexa488 (Figures 6B, S26, and S28)**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	gTCCCTgCCTCTATATCT	ΤT	TAgTAgTggTTTTACCggCgTCgAT	4	CACCggTgTAgAACAgAATACgTTC	TT	CCACTCAACTTTAACCCg
7	gTCCCTgCCTCTATATCT	ΤT	TggTAATACCACgTTCCTgCTCCTg	8	AgAATgCAgTAgTCgCAgCggAAgT	TT	CCACTCAACTTTAACCCg
11	gTCCCTgCCTCTATATCT	TT	CTTCgATTgTgAAgTCAACgTgCCC	12	CATCgAgAACACgCATggAACgTTC	TT	CCACTCAACTTTAACCCg
15	gTCCCTgCCTCTATATCT	TT	CTTTATATTTgTTTgCCTgACgCCA	16	TTTTgTTAACgAACgCAATgCgCgg	TT	CCACTCAACTTTAACCCg
19	gTCCCTgCCTCTATATCT	TT	gCTgCAgCggAACCgggTTCgCgCC	20	TgAAATgTTCTTCAgCACCAATCgC	TT	CCACTCAACTTTAACCCg
23	gTCCCTgCCTCTATATCT	ΤT	TATCTTCgTATTCgAAggTTACgCC	24	TAgCCAgTTCAACCATgTCTgCCgg	TT	CCACTCAACTTTAACCCg
27	gTCCCTgCCTCTATATCT	TT	gTTCTTCACCACCCAggTATTTTTC	28	gAgCACCTTTgATTTCTgCTTCAgT	TT	CCACTCAACTTTAACCCg
31	gTCCCTgCCTCTATATCT	TT	gCATCgCCTgAACACCTTTgTTCTT	32	ATggCAggTAATCAATTACCgCATC	TT	CCACTCAACTTTAACCCg
35	gTCCCTgCCTCTATATCT	TT	CgTCATCACTTgCgTgACgTTCAgC	36	TTTTgAACgCCAgTgCAgAgAACgg	TT	CCACTCAACTTTAACCCg
39	gTCCCTgCCTCTATATCT	TT	TCAgTACggTATCACCAgAgTTAAC	40	AACgCTCACgTgCAgCTTTCACggA	TT	CCACTCAACTTTAACCCg
43	gTCCCTgCCTCTATATCT	TT	TgATCggCgCATCCgggTCACACAg	44	gCTCAgggAATTCCATACgTTCCAg	TT	CCACTCAACTTTAACCCg
47	gTCCCTgCCTCTATATCT	TT	CTTTAgCCAgACggCCCAgAgCCAg	48	CAgTCCATACACggAAAgACgggTC	TT	CCACTCAACTTTAACCCg
51	gTCCCTgCCTCTATATCT	TT	ggATAgTTTCACggTAAgCAACCTg	52	TACCTTCAACATCggTAACTTTCTg	TT	CCACTCAACTTTAACCCg
55	gTCCCTgCCTCTATATCT	TT	ggTTTgAACCCggCTCCAgCgggTA	56	TgTCgTTgATgAACTCgTAgCCTTT	TT	CCACTCAACTTTAACCCg
59	gTCCCTgCCTCTATATCT	TT	CCAgCggACCTgCTTTCAgCTgTTC	60	TACCCATgTCTACTACCgggTAgCC	TT	CCACTCAACTTTAACCCg
63	gTCCCTgCCTCTATATCT	TT	CTTTAAAggCgATAgAAgCAgCCAg	64	gAACTggTTTCgCTTTCTTAAAgCC	TT	CCACTCAACTTTAACCCg
67	gTCCCTgCCTCTATATCT	TT	gACgACggCTCAAgTCACCgATAAC	68	CAgATTCCTgACCTTTgAgCATACC	TT	CCACTCAACTTTAACCCg
71	gTCCCTgCCTCTATATCT	ΤT	TggTCAgAgAACgCAgCTgAgTTgC	72	ATTCCATAgTgTATgATgCACgACC	TT	CCACTCAACTTTAACCCg

#### Organism: E. coli

### Target mRNA: isocitrate dehydrogenase (*icd*) Probe set: 20 split-initiator probe pairs (each probe carries half an HCR initiator) HCR amplifier: B1-Alexa594 (Figure S28)

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gAggAgggCAgCAAACgg	AA	CCggAACAACTACTTTACTTTCCAT	2	TTTgCAgggTgATCTTCTTgCCTTg	TA	gAAgAgTCTTCCTTTACg
3	gAggAgggCAgCAAACgg	AA	gATTTTCAggAACgTTgAgTTTgCC	4	CATCACCTTCAATgTAAgggATAAT	TA	gAAgAgTCTTCCTTTACg
5	gAggAgggCAgCAAACgg	AA	TggCTggggTTACATCTACACCgAT	6	CgACTgCAgCgTCgACCACTTTCAg	TA	gAAgAgTCTTCCTTTACg
7	gAggAgggCAgCAAACgg	AA	TTTTACgCTCgCCTTTATAggCTTT	8	CACCggTgTAAATTTCCATCCAggA	TA	gAAgAgTCTTCCTTTACg
9	gAggAgggCAgCAAACgg	AA	ACCTGACCATAAACCTGTGTGGATTT	10	CAAgAgTTTCAgCAggCAgCCAgAC	TA	gAAgAgTCTTCCTTTACg
11	gAggAgggCAgCAAACgg	AA	TggCAACgCgATATTCACgAATCAg	12	CAACCggAgTggTCAgCggACCTTT	TA	gAAgAgTCTTCCTTTACg
13	gAggAgggCAgCAAACgg	AA	CAACgTTCAgAgAgCgAATACCgCC	14	TgTAgAgATCCAgTTCCTggCgCAg	TA	gAAgAgTCTTCCTTTACg
15	gAggAgggCAgCAAACgg	AA	gATAgTAACgTACCggACgCAggCA	16	ggTgTTTAACCggggCTTggAgTgCC	TA	gAAgAgTCTTCCTTTACg
17	gAggAgggCAgCAAACgg	AA	ggAAgATAACCATATCggTCAgTTC	18	CCgCATAAATgTCTTCCgAgTTTTC	TA	gAAgAgTCTTCCTTTACg
19	gAggAgggCAgCAAACgg	AA	CggCAgAgTCTgCTTTCCATTCgAT	20	gCAggAATTTAATCACTTTCTCggC	TA	gAAgAgTCTTCCTTTACg
21	gAggAgggCAgCAAACgg	AA	gAATTTTCTTCACCCCATCTCTTC	22	TACCgATACCACAATgTTCCgggAA	TA	gAAgAgTCTTCCTTTACg
23	gAggAgggCAgCAAACgg	AA	TggTgCCTTCTTCCgAACACggCTT	24	ATTCgATCgCTgCACgAACCAgACg	TA	gAAgAgTCTTCCTTTACg
25	gAggAgggCAgCAAACgg	AA	CAgAgTCACgATCgTTAgCAATTgC	26	TgATgTTgCCTTTgTgCACCAgAgT	TA	gAAgAgTCTTCCTTTACg
27	gAggAgggCAgCAAACgg	AA	CTTTAAACgCTCCTTCggTgAACTT	28	CTTCACgCgCCAgCTggTAgCCCCA	TA	gAAgAgTCTTCCTTTACg
29	gAggAgggCAgCAAACgg	AA	CACCgTCgATCAgTTCACCgCCAAA	30	TCgggTTTTTAACTTTCAgCCACgg	TA	gAAgAgTCTTCCTTTACg
31	gAggAgggCAgCAAACgg	AA	CTTTAATgACgATCTCTTTgCCAgT	32	gTTgCAggAATgCATCAgCAATCAC	TA	gAAgAgTCTTCCTTTACg
33	gAggAgggCAgCAAACgg	AA	CATATTCAgCCggACgCAgCAggAT	34	CgTTCAggTTCATACAggCgATAAC	TA	gAAgAgTCTTCCTTTACg
35	gAggAgggCAgCAAACgg	AA	CTgCCAgggCgTCAgAAATgTAgTC	36	gggCgATACCgATACCgCCAACCTg	TA	gAAgAgTCTTCCTTTACg
37	gAggAgggCAgCAAACgg	AA	CgCATTCgTCACCgATgTTTgCACC	38	CAgTACCgTgggTggCTTCAAACAg	TA	gAAgAgTCTTCCTTTACg
39	gAggAgggCAgCAAACgg	AA	CTTTgTCCTgACCggCATATTTCgg	40	CggAgAgAATAATAgAgCCAggATT	TA	gAAgAgTCTTCCTTTACg

## S4.9 Split-initiator probes for Figures 7, S29, and S30

Organism: H. sapiens sapiens

Target mRNA: **B-Raf proto-oncogene, serine/threonine kinase** (*BRAF*) Probe set: **23 split-initiator probe pairs** (each probe carries half an HCR initiator) HCR amplifier: **B3-Alexa647** 

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gTCCCTgCCTCTATATCT	TT	ACCgCTCAgCgCCgCCATCTTATAA	2	gCCCggCTCCgCgCCgCCACCACCg	ΤT	CCACTCAACTTTAACCCg
5	gTCCCTgCCTCTATATCT	TT	ggCAgggTCCgCAgCCgAAgAggCC	6	TTTgATATTCCACACCTCCTCCggA	ΤT	CCACTCAACTTTAACCCg
9	gTCCCTgCCTCTATATCT	TT	ATATATTgATggTggATTATgCTCC	10	gCTggTgTATTCTTCATAggCCTCC	ΤT	CCACTCAACTTTAACCCg
13	gTCCCTgCCTCTATATCT	TT	AgAgCTAgAAACAgAAAAATCAgTT	14	AgAAgATgTAACggTATCCATTgAT	ΤT	CCACTCAACTTTAACCCg
17	gTCCCTgCCTCTATATCT	TT	CTTggggTTgCTCCgTgCCACATCT	18	gACTCTAACgATAggTTTTTgTggT	TT	CCACTCAACTTTAACCCg
21	gTCCCTgCCTCTATATCT	TT	CTCTCCATCCTgAATTCTgTAAACA	22	ATCAgTgTCCCAACCAATTggTTTC	ΤT	CCACTCAACTTTAACCCg
25	gTCCCTgCCTCTATATCT	TT	TTTTCgTACAAAgTTgTgTgTTgTA	26	gTCACAAAATgCTAAggTgAAAAAC	ΤT	CCACTCAACTTTAACCCg
29	gTCCCTgCCTCTATATCT	TT	AACTTCTgTACTACAACgCTggTgA	30	TTggTCATAATTAACACACATCAgT	TT	CCACTCAACTTTAACCCg
33	gTCCCTgCCTCTATATCT	TT	TAgggCAgTCTCTgCTAAggACgCC	34	gggTgCggAAggggATgATCCAgAT	TT	CCACTCAACTTTAACCCg
37	gTCCCTgCCTCTATATCT	TT	TggTCggAAgggCTgTggAATTggA	38	AAATTGATTTCGATGATCTTCATCT	ΤT	CCACTCAACTTTAACCCg
41	gTCCCTgCCTCTATATCT	TT	TCTAATCAAgTCATCAATATTgACA	42	TCCTCCATCACCACgAAATCCTTgg	ΤT	CCACTCAACTTTAACCCg
45	gTCCCTgCCTCTATATCT	TT	TggAgATTTCTgTAAggCTTTCACg	46	AgATgACTTCCTTTCTCgCTgAggT	ΤT	CCACTCAACTTTAACCCg
49	gTCCCTgCCTCTATATCT	TT	CTgCCCATCAggAATCTCCCAATCA	50	AgATCCAATTCTTTgTCCCACTgTA	ΤT	CCACTCAACTTTAACCCg
53	gTCCCTgCCTCTATATCT	TT	AggTgTAggTgCTgTCACATTCAAC	54	TTCATTTTTgAAggCTTgTAACTgC	ΤT	CCACTCAACTTTAACCCg
57	gTCCCTgCCTCTATATCT	TT	ggTCTCAATgATATggAgATggTgA	58	ATCTATAAgTTTgATCATCTCAAAT	ΤT	CCACTCAACTTTAACCCg
61	gTCCCTgCCTCTATATCT	TT	CACTGTAGCTAGACCAAAATCACCT	62	CTgATgggACCCACTCCATCgAgAT	TT	CCACTCAACTTTAACCCg
65	gTCCCTgCCTCTATATCT	TT	CTgAAAgCTgTATggATTTTTATCT	66	AACAATTCCAAATgCATATACATCT	TT	CCACTCAACTTTAACCCg
69	gTCCCTgCCTCTATATCT	TT	ACTAAAATCCTCTgTTTggAAACCA	70	TgTTTTTggAgAAgCACAAgCATAT	TT	CCACTCAACTTTAACCCg
73	gTCCCTgCCTCTATATCT	TT	TTTTgTTgCTACTCTCCTgAACTCT	74	AAgCAAACATATgTTCATTTATTTT	TT	CCACTCAACTTTAACCCg
77	gTCCCTgCCTCTATATCT	TT	ATTATATCTAgTCTTTAACCACACA	78	ТААдТАТАААТТТТАдТТТддддАА	TT	CCACTCAACTTTAACCCg
81	gTCCCTgCCTCTATATCT	TT	AAgTAAAgCCTCTAgAAgAggCTCT	82	ААдТдААТдАТАСАААСССддААСА	TT	CCACTCAACTTTAACCCg
85	gTCCCTgCCTCTATATCT	TT	TCTTCTggAgTCCCTAgTggACATg	86	CTgCAAACACAggCATAggTAgggT	ΤT	CCACTCAACTTTAACCCg
89	gTCCCTgCCTCTATATCT	TT	ATTAAATTCTACTgACTTCCTAAAT	90	ΑΑΑΑΤΤΑΤΤΑΑΘΑΑΤΑΑΤΑΑΤΑΘΑΑ	TT	CCACTCAACTTTAACCCg
## Organism: *H. sapiens sapiens* Target mRNA: **B-Raf proto-oncogene, serine/threonine kinase** (*BRAF*) Probe set: **23 split-initiator probe pairs** (each probe carries half an HCR initiator) HCR amplifier: **B4-Alexa546**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	CCTCAACCTACCTCCAAC	AA	CTCCATgTCCCCgTTgAACAgAgCC	4	ggCgCCggCgCCggCgCCggCCTCg	AT	TCTCACCATATTCgCTTC
7	CCTCAACCTACCTCCAAC	AA	ATgTTCCTgTgTCAACTTAATCATT	8	ACCAAATTTgTCCAATAgggCCTCT	AT	TCTCACCATATTCgCTTC
11	CCTCAACCTACCTCCAAC	AA	TTCTCTTTgTTggAgTgCATCTAgC	12	gTTCCCCAgAgATTCCAATAACTgT	AT	TCTCACCATATTCgCTTC
15	CCTCAACCTACCTCCAAC	AA	AggTAgCACTgAAAggCTAgAAgAg	16	gggATTTTgAAAAACTgAAAgAgAT	AT	TCTCACCATATTCgCTTC
19	CCTCAACCTACCTCCAAC	AA	CACTgTCCTCTgTTTgTTgggCAgg	20	gACTgTAACTCCACACCTTgCAggT	AT	TCTCACCATATTCgCTTC
23	CCTCAACCTACCTCCAAC	AA	CAATTCTTCTCCAgTAAgCCAggAA	24	TggAACATTCTCCAACACTTCCACA	AT	TCTCACCATATTCgCTTC
27	CCTCAACCTACCTCCAAC	AA	ACCCTggAAAAgCAgCTTTCgACAA	28	TTTATAACCACATgTTTgACAgCgg	AT	TCTCACCATATTCgCTTC
31	CCTCAACCTACCTCCAAC	AA	gAACTTggAgACAAACAgCAAATCA	32	TTCCTgTggTATTgggTggTgTTCA	AT	TCTCACCATATTCgCTTC
35	CCTCAACCTACCTCCAAC	AA	AATTTggggCCCAATAgAgTCCgAg	36	ggATTTTgAAggAgACggACTggTg	AT	TCTCACCATATTCgCTTC
39	CCTCAACCTACCTCCAAC	AA	AgCTgATgAggATCggTCTCgTTgC	40	TTCTATTgTgTTTATATgCACATTg	AT	TCTCACCATATTCgCTTC
43	CCTCAACCTACCTCCAAC	AA	gggggTAgCAgACAAACCTgTggTT	44	AgTTAgTgAgCCAggTAATgAggCA	AT	TCTCACCATATTCgCTTC
47	CCTCAACCTACCTCCAAC	AA	CATTCgATTCCTgTCTTCTgAggAT	48	ACTCgAgTCCCgTCTACCAAgTgTT	AT	TCTCACCATATTCgCTTC
51	CCTCAACCTACCTCCAAC	AA	TCCCTTgTAgACTgTTCCAAATgAT	52	TTTCACTgCCACATCACCATgCCAC	AT	TCTCACCATATTCgCTTC
55	CCTCAACCTACCTCCAAC	AA	ATgTCgTgTTTTCCTgAgTACTCCT	56	ATAgCCCATgAAgAgTAggATATTC	AT	TCTCACCATATTCgCTTC
59	CCTCAACCTACCTCCAAC	AA	TATATTATTACTCTTgAggTCTCTg	60	TTTTACTgTgAggTCTTCATgAAgA	AT	TCTCACCATATTCgCTTC
63	CCTCAACCTACCTCCAAC	AA	CAAAATggATCCAgACAACTgTTCA	64	CATTCTgATgACTTCTggTgCCATC	AT	TCTCACCATATTCgCTTC
67	CCTCAACCTACCTCCAAC	AA	CTTCATggCTTTTggACAgTTACTC	68	CTTTTTgAggCACTCTgCCATTAAT	AT	TCTCACCATATTCgCTTC
71	CCTCAACCTACCTCCAAC	AA	CgCACCATATCCCCCTgCCTggATg	72	CACTCATTTgTTTCAgTggACAggA	AT	TCTCACCATATTCgCTTC
75	CCTCAACCTACCTCCAAC	AA	gAgAgTATTTTATTCAATTTAACAT	76	TgTTCTTTggTTCACCTTAAAAAAA	AT	TCTCACCATATTCgCTTC
79	CCTCAACCTACCTCCAAC	AA	AACCCTTggATgTTAAAAATCCAAT	80	CAATTTTTAgCAATgTCTATgTATT	AT	TCTCACCATATTCgCTTC
83	CCTCAACCTACCTCCAAC	AA	AACTGAAGTTTACTACTTAAAATAA	84	ATAgCTggCAACAAAAgTTgCATgA	AT	TCTCACCATATTCgCTTC
87	CCTCAACCTACCTCCAAC	AA	CAggCTAACCgACTgCCAACTTCTC	88	gATCTgTTCAgTTTgCCTTATCTAA	AT	TCTCACCATATTCgCTTC
91	CCTCAACCTACCTCCAAC	AA	ΑΤΤ <u></u> GTTATAAAAAgAAATAgTTATA	92	gAAATAAAAgACATCCACATTTTCC	AT	TCTCACCATATTCgCTTC

## Organism: *G. gallus domesticus* Target mRNA: **diencephalon/mesencephalon homeobox 1** (*Dmbx1*) Probe set: **25 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B1-Alexa594**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
1	gAggAgggCAgCAAACgg	AA	CgCTCAgTgAgTTCATggCATgCAg	2	CTgCCTgCTggTgCAggTTgTACAT	TA	gAAgAgTCTTCCTTTACg
5	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	6	ggTgCTgggATCCATAACgTgCCTC	TA	gAAgAgTCTTCCTTTACg
9	gAggAgggCAgCAAACgg	AA	AAATgATgTCAgCCAgTCTCTCCgC	10	gCTTCTgCAgCTgCTCCTTCTggAg	TA	gAAgAgTCTTCCTTTACg
13	gAggAgggCAgCAAACgg	AA	CAgTgTCAggTATTgTggACTgggT	14	CCTCACTTgggATgCTCTgAgTCTT	TA	gAAgAgTCTTCCTTTACg
17	gAggAgggCAgCAAACgg	AA	CCTCCTCTCTgTCAgTCTggTCCTC	18	TAgCCTCATCCAAggTgCTCTTAAA	TA	gAAgAgTCTTCCTTTACg
21	gAggAgggCAgCAAACgg	AA	CgCTgATgggAgACTCTgATTTTgg	22	CACTgCTAgATgAAggAgTCACggT	TA	gAAgAgTCTTCCTTTACg
25	gAggAgggCAgCAAACgg	AA	CTgCCATgTgCTggCggAATTgCTC	26	AggAgTAATggACCAAgTTgTTggT	TA	gAAgAgTCTTCCTTTACg
29	gAggAgggCAgCAAACgg	AA	gACAgTgCAgggAgCTCAgAggCgC	30	CgTgggAgAgAgATTggTAgTACgA	TA	gAAgAgTCTTCCTTTACg
33	gAggAgggCAgCAAACgg	AA	TCTCAATACTTgTTgTTTTACTgTT	34	CATgCTgCTTTgCCCggAgCCTTAg	TA	gAAgAgTCTTCCTTTACg
37	gAggAgggCAgCAAACgg	AA	gACCTCCggTgCATCTTCTTATggg	38	ggTTCCAggAgTgACATgTCTggTg	TA	gAAgAgTCTTCCTTTACg
41	gAggAgggCAgCAAACgg	AA	CggTgCTAgTAAgACATTAgTAAAT	42	AgCCAgTAgCAgTgTCTgATgCAAT	TA	gAAgAgTCTTCCTTTACg
45	gAggAgggCAgCAAACgg	AA	TCACAgCAgTCCAAAgggACAgTTC	46	gCTCTTTCTgAATgTTACAggCTTA	TA	gAAgAgTCTTCCTTTACg
49	gAggAgggCAgCAAACgg	AA	gTTTTCCCAAAgAATgCATCgACAA	50	TATgTACAAgACAAAgCAggACTCT	TA	gAAgAgTCTTCCTTTACg
53	gAggAgggCAgCAAACgg	AA	ggggAATAAAAgCAAAAgAggCCAC	54	gACTAgCTACCAAAACTgAgAgAgA	TA	gAAgAgTCTTCCTTTACg
57	gAggAgggCAgCAAACgg	AA	TTTgCTCTAAgCACCATTAAgACTC	58	gAgCAgTgAATTgCATAATggTTTT	TA	gAAgAgTCTTCCTTTACg
61	gAggAgggCAgCAAACgg	AA	ggAAgTgCTTAAACAggAAATTCAC	62	AgTAAAggAAAAAACACTTgCCTTT	TA	gAAgAgTCTTCCTTTACg
65	gAggAgggCAgCAAACgg	AA	AATTTggCTTTCATTTTTCTCCCCA	66	AACAATCAAgTCAAAAgTAACCATg	TA	gAAgAgTCTTCCTTTACg
69	gAggAgggCAgCAAACgg	AA	CTAgACCAAAATgCTCTCCAAAAAg	70	AgTTTTTATTgTTCTCTATTgTCgA	TA	gAAgAgTCTTCCTTTACg
73	gAggAgggCAgCAAACgg	AA	TAAgAACAgCTTgCATTAATCgTgg	74	TAgAATTTggTgATCggAgCgTTTT	TA	gAAgAgTCTTCCTTTACg
77	gAggAgggCAgCAAACgg	AA	CTTggCCTCCAgCATTgCAgCATTT	78	AATAgAAAgCCCCCgATTATCACCC	TA	gAAgAgTCTTCCTTTACg
81	gAggAgggCAgCAAACgg	AA	gAAATCACTTTgCAgTTggTgAgTT	82	AgAgAAATgAggCCAAAACTgTggA	TA	gAAgAgTCTTCCTTTACg
85	gAggAgggCAgCAAACgg	AA	AAgTCCTTTgggTTggTAgAAAAgT	86	TATTTggTTTggAgAAgATAAATAA	TA	gAAgAgTCTTCCTTTACg
89	gAggAgggCAgCAAACgg	AA	gCATTTTTggTCCTAggCAATACTA	90	ACCTAgCACCTgCCACAgAgCCAgT	TA	gAAgAgTCTTCCTTTACg
93	gAggAgggCAgCAAACgg	AA	gACCAgATgATggCCTgCAgTgAAT	94	CTCCATTTCTTCTTTAAATCgAgCA	TA	gAAgAgTCTTCCTTTACg
97	gAggAgggCAgCAAACgg	AA	AggTAACTCAACCCACggCTTCTgC	98	AACACCCTTTCCCCCTCgTgTTTAA	TA	gAAgAgTCTTCCTTTACg

## Organism: *G. gallus domesticus* Target mRNA: **diencephalon/mesencephalon homeobox 1** (*Dmbx1*) Probe set: **25 split-initiator probe pairs (each probe carries half an HCR initiator)** HCR amplifier: **B2-Alexa647**

Odd #	1st Half of Initiator I1	Spacer	Probe Sequence (25 nt)	Even #	Probe Sequence (25 nt)	Spacer	2nd Half of Initiator I1
3	CCTCgTAAATCCTCATCA	AA	AgTCgggCgCgTgCTgTgCTTgCTg	4	gggTgAgggCgTgCACTgAgggCCg	AA	ATCATCCAgTAAACCgCC
7	CCTCgTAAATCCTCATCA	AA	AggCCgTgCgACTTCgCCTTTgTTT	8	CCAgTgCCTCCAgCTgTTgggCAgT	AA	ATCATCCAgTAAACCgCC
11	CCTCgTAAATCCTCATCA	AA	CACTgTggCTgCTTTCACAgTCCTT	12	CCAgCACAggTggTTCCgTCTTCCC	AA	ATCATCCAgTAAACCgCC
15	CCTCgTAAATCCTCATCA	AA	ACAgggTTAggTTgAggTCTgTgCT	16	gCgCTgATTCACTggCTgACTgCTC	AA	ATCATCCAgTAAACCgCC
19	CCTCgTAAATCCTCATCA	AA	CgTCCACACCAggACTCTTgTCCAC	20	TCgCTCTCTTgCAgTTCAAAgCCTT	AA	ATCATCCAgTAAACCgCC
23	CCTCgTAAATCCTCATCA	AA	AggAgTAggAgTgAgTTTgAgCCAg	24	gCAggCggAAgAggCTCAggggTgA	AA	ATCATCCAgTAAACCgCC
27	CCTCgTAAATCCTCATCA	AA	TggCAgACggTgTCCCCATTTCgAA	28	TgTTgACATTCATgCCCAAgTAggg	AA	ATCATCCAgTAAACCgCC
31	CCTCgTAAATCCTCATCA	AA	ggATgggTgAggACCAgACCTgCTg	32	ggCTTggAAgAgAgCTggAggCCTg	AA	ATCATCCAgTAAACCgCC
35	CCTCgTAAATCCTCATCA	AA	gTAgggTgTCAAgCCCAAgggATgC	36	ACggTTgCCCATCTggCAgTCAgTT	AA	ATCATCCAgTAAACCgCC
39	CCTCgTAAATCCTCATCA	AA	ACTCCAggAAgAgATgAgggTggAA	40	AAAgTTTTCCCTgATAgggAgCACC	AA	ATCATCCAgTAAACCgCC
43	CCTCgTAAATCCTCATCA	AA	TCCTCCTCAAATATTTAAAgAAgAC	44	CTgTCTAAACACACATCCTCTCCCT	AA	ATCATCCAgTAAACCgCC
47	CCTCgTAAATCCTCATCA	AA	ACATTATCgCAgggATgAggTgAgg	48	AAAAAgggTgTATATAACACggTTg	AA	ATCATCCAgTAAACCgCC
51	CCTCgTAAATCCTCATCA	AA	gCggTggATgCTTTCAACATTgTAA	52	TTCTgTAACACTgACAgTAACACAC	AA	ATCATCCAgTAAACCgCC
55	CCTCgTAAATCCTCATCA	AA	gggAgCgTggCTgATTTgTgACTTT	56	AACCCAAgAAgAgCAACTAgCTgTg	AA	ATCATCCAgTAAACCgCC
59	CCTCgTAAATCCTCATCA	AA	TCAgCTTTAgCAgAgAAgAgAgAAg	60	TTgCATCATTTCCTggCCgTTATAA	AA	ATCATCCAgTAAACCgCC
63	CCTCgTAAATCCTCATCA	AA	TCTgTCTgTgAACAAgTgCTATTAg	64	CAgCAgCATTTggCCAgCATTTTgT	AA	ATCATCCAgTAAACCgCC
67	CCTCgTAAATCCTCATCA	AA	TTACACTTCACTgAAgACCAAAgAg	68	AACCCCATAATTTgTAAATgggggA	AA	ATCATCCAgTAAACCgCC
71	CCTCgTAAATCCTCATCA	AA	gTATgAACACAgTgggAgTTCATAC	72	TTgTCAAggAACCATATAATTCATg	AA	ATCATCCAgTAAACCgCC
75	CCTCgTAAATCCTCATCA	AA	AgggTCTgAAgCTgCACAgCTTgAg	76	CACTTGTTACATTCTCACTTGCTAA	AA	ATCATCCAgTAAACCgCC
79	CCTCgTAAATCCTCATCA	AA	CAAgCCAATCTACTCCTCgCTgCAg	80	ggTTgCTTggggACATggTACTTTT	AA	ATCATCCAgTAAACCgCC
83	CCTCgTAAATCCTCATCA	AA	TggATTACTAAAATgAAgggTCATT	84	CTTCTCAAgAAggAAAAACACTCTg	AA	ATCATCCAgTAAACCgCC
87	CCTCgTAAATCCTCATCA	AA	TTCTTggTACggTgAgTTCAAAggA	88	TTAgATCTgggTTTCCTCCCTCCCT	AA	ATCATCCAgTAAACCgCC
91	CCTCgTAAATCCTCATCA	AA	gACCCgCCTgACACCCTTTggATTC	92	gCCCAgCTCTgCTgCgTgTTAgTgg	AA	ATCATCCAgTAAACCgCC
95	CCTCgTAAATCCTCATCA	AA	gTTATgTAggCTATgCACACgTTgC	96	TggTATgAAgTAAgATgggAgCAAg	AA	ATCATCCAgTAAACCgCC
99	CCTCgTAAATCCTCATCA	AA	TTTTTAAgATgCATTATTgCAgTTg	100	TgCCTCAgTTTAAgggATTTAgATg	AA	ATCATCCAgTAAACCgCC

## References

- Bell, G.W., Yatskievych, T.A., & Antin, P.B. (2004). GEISHA, a whole-mount in situ hybridization gene expression screen in chicken embryos. *Dev. Dyn.*, 229(3), 677–687.
- Choi, H. M. T., Chang, J. Y., Trinh, L. A., Padilla, J. E., Fraser, S. E., & Pierce, N. A. (2010). Programmable in situ amplification for multiplexed imaging of mRNA expression. *Nat. Biotechnol.*, **28**(11), 1208–12.
- Choi, H. M. T., Beck, V. A., & Pierce, N. A. (2014). Next-generation in situ hybridization chain reaction: higher gain, lower cost, greater durability. *ACS Nano*, **8**(5), 4284–4294.
- Choi, H. M. T., Calvert, C. R., Husain, N., Huss, D., Barsi, J. C., Deverman, B. E., Hunter, R. C., Kato, M., Lee, S. M., Abelin, A. C. T., Rosenthal, A. Z., Akbari, O. S., Li, Y., Hay, B. A., Sternberg, P. W., Patterson, P. H., Davidson, E. H., Mazmanian, S. K., Prober, D. A., van de Rijn, M., Leadbetter, J. R., Newman, D. K., Readhead, C., Bronner, M. E., Wold, B., Lansford, R., Sauka-Spengler, T., Fraser, S. E., & Pierce, N. A. (2016). Mapping a multiplexed zoo of mRNA expression. *Development*, 143, 3632–3637.
- Couprie, M., & Bertrand, G. (1997). Topological gray-scale watershed transform. *Pages 136–146 of: Vision geometry v.* Proceedings of SPIE.
- Darnell, D. K., Kaur, S., Stanislaw, S., Davey, S., Konieczka, J. H., Yatskievych, T. A., & Antin, P. B. (2007). GEISHA: an in situ hybridization gene expression resource for the chicken embryo. *Cytogenet. Genome Res.*, 117(1-4), 30–35.
- McEntyre, J., & Ostell, J. (2002). *The NCBI handbook [Internet]*. Bethesda, MD: National Center for Biotechnology Information (US).
- Shah, S., Lubeck, E., Schwarzkopf, M., He, T.-F., Greenbaum, A., Sohn, C. H., Lignell, A., Choi, H. M. T., Gradinaru, V., Pierce, N. A., & Cai, L. (2016). Single-molecule RNA detection at depth via hybridization chain reaction and tissue hydrogel embedding and clearing. *Development*, 143, 2862–2867.
- Trivedi, V., Choi, H. M. T., Fraser, S. E., & Pierce, N. A. (2018). Multidimensional quantitative analysis of mRNA expression within intact vertebrate embryos. *Development*, **145**, DOI:10.1242/dev.156869.
- Yoo, T.S., Ackerman, M.J., Lorensen, W.E., Schroeder, W., Chalana, V., Aylward, S., Metaxas, D., & Whitaker, R. (2002). Engineering and algorithm design for an image processing API: a technical report on ITK - the Insight Toolkit. *Studies in Health Technology and Informatics*, 85, 586–592.