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The Asymmetric Cell Division Regulators Par3, Scribble and Pins/Gpsm2 Are Not Essential for Erythroid Development or Enucleation

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Abstract

Erythroid enucleation is the process by which the future red blood cell disposes of its nucleus prior to entering the blood stream. This key event during red blood cell development has been likened to an asymmetric cell division (ACD), by which the enucleating erythroblast divides into two very different daughter cells of alternate molecular composition, a nucleated cell that will be removed by associated macrophages, and the reticulocyte that will mature to the definitive erythrocyte. Here we investigated gene expression of members of the Par, Scribble and Pins/Gpsm2 asymmetric cell division complexes in erythroid cells, and functionally tested their role in erythroid enucleation *in vivo* and *ex vivo*. Despite their roles in regulating ACD in other contexts, we found that these polarity regulators are not essential for erythroid enucleation, nor for erythroid development *in vivo*. Together our results put into question a role for cell polarity and asymmetric cell division in erythroid enucleation.

Introduction

Erythroid enucleation embodies many features of an asymmetric cell division [1-10] and results in the generation of two unique daughter cells: the pyrenocyte containing the condensed nucleus and the anucleated reticulocyte that will further mature into the erythrocyte found in the peripheral blood. One of the first visible steps of establishing polarity prior to nuclear extrusion is the phosphoinositide 3-kinase/microtubule -directed migration of the nucleus to one side of the cytoplasm, whereby the nucleus apposes the cell membrane [5]. This



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event is followed by a polarized enrichment of cytoskeletal elements, including actin, and myosin II to the future restriction site between pyrenocyte and future red blood cell[6–9]. During the enucleation process specific membrane proteins are either sorted to the nucleus (e.g. Emp protein) or to the future reticulocyte (e.g. spectrin)[7, 10].

Asymmetric cell division (ACD) is crucial for generating cell progeny with diverse functions and requires a polarized distribution of cell fate determinants, cytoskeletal regulators and polarity proteins. The pathways regulating asymmetric cell division have mainly been derived from studies in Drosophila melanogaster, particularly from neuroblasts, which undergo up to 20 rounds of asymmetric cell divisions to generate neurons of the larval nervous system[11]. Three cell polarity complexes have been identified to contribute to the establishment of asymmetry in *Drosophila* neuroblasts: the Par, Pins and Scribble complexes^[12]. The Pins complex (comprised of Inscuteable (Insc), Partner of Insc (Pins) and a Gai subunit) regulates mitotic spindle orientation by providing an attachment site for astral microtubules^[13]. The Par complex (made up of Par3, Par6 and atypical Protein Kinase C (aPKC)) and the Scribble complex (comprised of Scribble, Discs Large (Dlg) and Lethal Giant Larvae (Lgl)) are required for the establishment of cell polarity that is critical for ACD[14]. All of these complexes are conserved in vertebrates and are thought to play similar roles. For example, Par3 has been implicated in determining polarity in mammalian oogenesis[15], an example of an asymmetric cell division, in which the egg divides to produce an oocyte and a small polar body. Cells of the hematopoietic systems make fate decisions in order to either self-renew, differentiate, proliferate or to undergo apoptosis. ACD may control these decisions[16]. For example, it has been suggested that T-cell development displays many features characteristic of ACD[17]. In line with these observations, studies using knockdown and knockout approaches implicated important roles for the Scribble and Par3 complexes in the development and function of lymphocytes[18]. Pins, also known as Gpsm2 (G protein signaling modulator 2) is thought to enhance haematopoietic stem cell function through altered asymmetric and symmetric divisions[19].

Despite the proposal that erythroid enucleation embodies many features of ACD, the requirement for ACD regulators in this event has not been investigated to date. Given that ACD utilizes a conserved molecular toolbox across species and within different forms of tissue development[20], we examined expression of known critical asymmetric division genes, from the Scribble, Par and Pins complex in orthochromatic erythroblasts, and used the corresponding mouse models to functionally test the role of ACD regulators in erythroid development and enucleation.

Materials and Methods

Materials

Phenylhydrazine hydrochloride (PHZ) was purchased from Aldrich Chemistry. PE-Cy7 conjugated anti-CD44 mouse antibodies were purchased from BD Pharmingen. Alexa Fluor 647 conjugated anti-Ter119 mouse antibodies were purchased from Biolegend. Hoechst 33342 was purchased from Invitrogen. Propidium iodide (PI) was purchased from Merck. Rapid Diff stain was purchased from Australian Biostain.

Animal experiments

This study was carried out in strict accordance with the recommendations of the Victorian Bureau of Animal Welfare, Department of Primary Industries, and the National Health and Medical Research Council's Australian code of practice for the care and use of animals for scientific purposes. The protocol was approved by the Institutional Animal Care and Use Committee: Peter MacCallum Cancer Centre Animal Experimentation Ethics Committee under Permit number E535. All efforts were made to minimize suffering. All mice (71 females, 72 males) used in this study were on a C57BL/6 background and 6–12 weeks of age. Mice were kept at 21°C, with a humidity of approximately 60% on a 14h light/10h dark cycle and fed with standard mouse cubes (Ridley Agri). EpoR-Cre^{ki/+} mice[21] (referred to as *EpoR-Cre⁺* in this manuscript) were provided by Dr. Carl Walkley (St. Vincent's Institute, Melbourne, Australia). Conditional Par3 knockout mice (*Pard3*^{fl}) were generated by the laboratory of Prof. Shigeo Ohno (Yokohama City University, Yokohama, Japan)[22]. Conditional Scrib knockout mouse allele (*Scrib*^{fl}) were generated in-house[23]. Gpsm2^{Δ C} mice were generated by Fumio Matsuzaki[24]. To induce stress erythropoiesis mice at 6–12 weeks of age were administered intraperitoneal injections of phenylhydrazine hydrochloride (60µg/g) on day 0 and day 1 of the experiments. On day 4, cells were isolated from bone marrow and spleens. Following phenylhydrazine hydrochloride treatment, mice were monitored for signs of discomfort every two hours as defined by lethargy, ruffled fur or a hunched appearance, at which time the mice were considered to have reached the ethically permitted humane endpoint criteria and were humanely euthanized using cervical dislocation.

FACS analysis of erythropoiesis and enucleation

For analysis of erythropoiesis cells were harvested from bone marrow or spleen and stained for Ter119 and CD44[25]. PI was used to exclude dead cells from the sort. Erythropoiesis was analyzed using FACS LSR II. 20000 viable cells were analyzed for each sample.

Analysis of enucleation was done as previously described[26]. Shortly, *ex vivo* cells isolated from bone marrow or spleen were stained for CD44, Ter119, Hoechst and PI. All Hoechst negative (enucleated) cells were excluded from the sort. Orthochromatic erythroblasts were isolated based on their Ter119 and CD44 expression by FACS Aria II special order system (BD) using the FACS Diva software (BD). Enucleation was analyzed 5h post sort using FACS LSR II. 5000 cells were analyzed per sample. Net percentage of enucleation was then derived by dividing the number of enucleated cells (Ter119⁺/Hoechst⁻) by the sum of enucleated cells and erythroblasts (Ter119⁺/Hoechst⁺), and by subsequently multiplying the quotient by 100.

RNA isolation, cDNA synthesis and qPCR

Total RNA was extracted from FACS sorted erythroid cells, and reversed transcribed with Superscript III (Invitrogen). Real-time PCR amplification of cDNA was done in triplicates in a

Table I.	Sequences	of primers	used for	QRIPCR.	

Target	Forward sequence (5'-3')	Reverse sequence (5'-3')
β <i>2M</i>	TTCACCCCCACTGAGACT	GTCTTGGGCTCGGCCATA
scribble	GAGGAGATTTACCGCTACAGTCGTA	TGTCACTGAGGCCCAACTTTC
lgl1	TGTTGTACGCTGCCTCTACTTTG	GGTGCCCGCCCACAT
Igl2	GGTGGATTCAACCAAAGCCAA	TTGGCTCTTTGAGTTCCTGACG
dlg1	GAGAGTGACGAAGTCGGAGTGA	AATCGGGCTCGTTCCTTCTT
dlg3	GGTGGAAAAGAAGAGCGAGCT	GCATCCTCTTGTCCTAGGT
dlg4	CCCAACATGGACTGTCTCTGTATAGT	GGGCGTGTCTTCATCTTGGT
pins	GGCCAGTTTCAGTAATTTCA	AGGCTAAAGAAGTCCTCATCCG
insc	GTTCGGCTCAGCTGTATGTCTC	GGTAAGTGCACACAGTGTGCCT
Арксζ	AGCCAACGGCCACCTCTT	CAGTACGCTCCCCTGTTAAAGC
Apkci	CAGGCTGTACGAGCTGAACAAG	GGAAATACATGAATCAAGAGTTCAGAATC
pard3	CCCAACTTTTCCCTCGATGATA	AACAACCCCAATGTTCTGCC
pard3 (to show KO)	GAGATATTCGGCAGTGAGCTG	GAAGGCGTGACCTCAATTTCA

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B	EpoR-Cre ⁺ ; Pard3 ^{+/+}	EpoR-Cre⁺; Pard3 ^{f/fl}	EpoR-Cre ⁺ ; Scrib ^{+/+}	EpoR-Cre ⁺ ; Scrib ^{fl/fl}	Gpsm2 +/+	Gpsm2 ^{∆C/∆C}
Red blood cell [x10 ¹² /L]	10.03 ± 0.47	9.93 ± 0.65	9.40 ± 0.43	9.32 ± 0.46	9.38 ± 0.43	9.18 ± 0.46
Hemoglobin [g/dL]	15.60 ± 0.48	15.0 ± 0.54	14.55 ± 0.65	14.70 ± 0.62	14.81 ± 0.37	14.76 ± 0.69
Hematocrit [L/L]	0.50 ± 0.03	0.48 ± 0.03	0.48 ± 0.02	0.47 ± 0.01	0.47 ± 0.02	0.46 ± 0.02
MCV [x10 ⁻¹⁵ /L]	49.78 ± 0.55	48.63 ± 1.23	50.35 ± 1.46	50.35 ± 1.65	49.60 ± 0.97	49.67 ± 1.23
RDW [%CV]	15.18 ± 0.95	16.17 ± 3.15	16.63 ± 1.63	15.67 ± 0.93	15.73 ± 1.66	14.96 ± 0.91



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Fig 1. The ACD regulators Par3, Scribble and Gpsm2 are not required for erythroid enucleation during steadystate erythropoiesis. (A) Representative images of blood smears from peripheral blood derived from age-matched mice of the indicated genotypes of the different mouse models. Scale bar = 10µm. (B) Whole blood analysis using the CELL-DYN Sapphire System was performed on peripheral blood derived from age-matched mice of the indicated genotypes of the indicated mouse models. Data represent the mean (+/- SD) of 3–7 independent mice. (C) Orthochromatic erythroblasts were enriched by FACS (Aria II) from bone marrow of mice of the indicated genotypes of the indicated mouse models and incubated in 96-well plates at 30.000 cells per well. Graphs showing enucleation efficiencies 5h post sort quantified by LSR II. Data represent the mean (+/- SD) of 3–4 independent experiments. *P< 0.05, **P< 0.01, ***P< 0.001, ***P< 0.0001 (unpaired student's t-test). (D) Whole blood derived from aged-matched mice of the indicated genotypes of the different mouse models was exposed to decreasing concentrations of sodium chloride (NaCI) and the degree of hemolysis measured colorimetrically. Data represent the mean (+/- SD) of 3–5 independent experiments.

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StepOnePlus Real-Time PCR system (Applied Biosystems, Carlsbad, CA) using the SYBER Green gene expression assay (Applied Biosystems, USA). All samples were normalized to β 2Macroglobulin (β 2M) control and fold change between samples was calculated using the comparative C(T) method. Primers used for qRT-PCR are listed below (Table 1).

Genotyping

Mice were genotyped from genomic DNA isolated from tail biopsies. To show deletion of the floxed allele, DNA was isolated from FACS sorted erythroid cells (Ter119 positive). *EpoR-Cre* WT/Tg alleles were detected using the following primers 5' - TCCCGGACCCCAAGTTTGAG-3', 5' - AAGCCTCTGCCCTGAGCATCAC-3' and 5' - GTGTGGGCTGCCCCTTCTGCCA-3' [27]. The PCR program employed was 95° C for 5 minutes, followed by 43 cycles of 95° C for 30seconds, 62° C for 30seconds, 72° C for 30seconds, and a final elongation step at 72° C for 5 minutes. *Pard3* WT/Floxed alleles were detected using the following primers 5' - AGGCTAGCCTGGGTGA TTTGAGACC-3' and 5' - TTCCCTGAGGCCTGACACTCCAGTC-3'. The PCR reaction was performed at 95° C for 5 minutes, followed by 35 cycles of 95° C for 30 seconds, 64° C for 30 seconds and 72° C for 30 seconds. A final elongation was performed at 72° C for 5 minutes. The recombined *Pard3* allele was detected using the following primers 5' - TACCGTTAACTGCA GCT CGGCTCTG-3' and 5' - AGCTGGCGCTGGTACCATCTCCTCC-3'. The PCR program employed was 94° C for 3 minutes, followed by 40 cycles of 94° C for 30 seconds, 64° C for 60 seconds, 72° C for 45 seconds. A final elongation was performed at 72° C for 30 minutes.

WT and LoxP-flanked *Scrib* alleles were detected using the following primer sequences 5' - GCCATGGTGGCAGAGGTTGG-3' and 5' - TGCTTTCTCCCAGACTCAGG-3'. The recombined *Scrib* allele was detected using the following primer sequence 5' - GAGAAAGTTGGGCCTCAG TG-3'. The PCR program employed was 95°C for 2 minutes, followed by 35 cycles of 94°C for 30 seconds, 56°C for 30 seconds, 72°C for 30 seconds, and an elongation step of 72°C for 10 minutes.

Gpsm2 WT and deleted *Gpsm2* alleles were detected using the following primer sequences 5′-TCCTCCATCTGCTGCCACTAAG-3′, 5′-ACAGCCACCGAAGGTCACAAAG-3′ and 5′-TTCAGTAGGTTACCACACCATCCTG-3′. The PCR program employed was 95°C for 5 minutes, followed by 43 cycles of 95°C for 30 seconds, 66°C for 30 seconds and 72°C for 1 minute, and an elongation step of 72°C for 5 minutes.

Osmotic fragility test

1µl of blood (from eye bleeds) was added to 200µl of various NaCl concentrations (1, 0.85, 0.75, 0.65, 0.6, 0.55, 0.5, 0.45, 0.4, 0.35, 0.3, 0.2, 0.1, 0% NaCl in dH₂O). The blood was gently mixed with the NaCl solution and incubated at room temperature for 30 minutes. The sample was gently mixed again and then spun in a microcentrifuge at 14.000rpm for 2 minutes. The supernatant was transferred into a well of a 96-well plate and absorption measured (OD 550nm) by a





Fig 2. The ACD regulators Par3, Scribble or Gpsm2 are not required for steady-state erythropoiesis in the bone marrow. (A) Graphs showing total cell numbers per femur (on left) and percentages of erythroid cells (Ter119 positive) in femurs (on right) isolated from age-matched



mice of the indicated genotypes of the different mouse models. Data are mean (+/- SD) of 3–6 independent experiments. **(B)** (i.) Diagram showing the distinct erythroid populations at the different maturation stages (I-V) that can be identified by FACS. (ii.) Representative FACS plot showing erythroid differentiation stages in the bone marrow. **(C)** Bar graphs showing percentages of erythroid cells at the different developmental stages (I-V) during homeostasis in the bone marrow harvested from age-matched mice of the indicated genotypes of the different mouse models. Data represent the mean (+/- SD) of 3–6 independent experiments. *P< 0.05, **P< 0.01, ***P< 0.001, ****P< 0.0001 (unpaired student's t-test).

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plate reader. Percent hemolysis was calculated by dividing the absorption measured for the sample by the absorption measured in the well containing that sample in 0% NaCl (max. hemolysis), and by subsequently multiplying the quotient by 100.

Results and Discussion

Late stage erythroblasts express polarity regulators

To test whether the core conserved ACD machinery was involved in erythroid enucleation, we first examined the expression of ACD regulators in terminal differentiating erythroblasts and found that orthochromatic erythroblasts (the stage prior to enucleation) express genes of the Scribble complex (*Scrib*, *Lgl1*, *Lgl2*, *Dlg1*, *Dlg3*, *Dlg4*), the Pins complex (*Gpsm2*, *Insc*), and the Par3 complex (*aPKCζ*, *aPKCi*, *Pard3*) (S1 Fig). Of all genes surveyed, *Pard3* was significantly upregulated in orthochromatic erythroblasts compared to earlier, proliferating stages (S1C Fig).

Characterization of knockout mouse models

We next tested if Par3, and other ACD regulators, Scribble and Gpsm2, play a role in erythroid enucleation. Par3 deficiency in the whole mouse results in defective cardiac development and embryonic lethality in the midgestational stage[22]. Similarly, Scribble null mice are neonatal lethal due to neural tube and abdominal wall closure defects[28, 29]. To investigate the biological functions of these polarity regulators in erythroid cells *in vivo*, mice harboring conditional *Pard3*[22] or *Scrib*[23] alleles were crossed to *Epor*-Cre^{ki/+} mice[21]. Expression of Cre recombinase from the endogenous *Epor* locus targets Cre expression to erythroid cells and therefore depletes Par3/Scribble specifically in the erythroid compartment as confirmed by PCR and qPCR analysis (S2A and S2B Fig). To assess Gpsm2 loss of function, we utilized *Gpsm2*^{AC} mice that carry a germline intragenic deletion of the GoLoco motif[24]. We confirmed the presence of this deletion by PCR and qPCR analysis (S2C Fig). Defective asymmetric cell divisions have been observed in these mice previously during neurogenesis[24].

Polarity regulators Par3, scribble and Gpsm2 are not essential for red blood cell development

Investigation of blood smears of peripheral blood collected from age-matched mice of the indicated genotypes revealed no enucleation defects in red blood cells in the absence of functional erythroid Par3, Scribble or Gpsm2 expression (Fig 1A (i-iii)). Whole blood analysis also confirmed no significant changes in other parameters including hemoglobin, hematocrit, mean corpuscular volume and red cell distribution width in mice with erythroid cells deficient for Par3, Scribble or Gpsm2 (Fig 1B (i-iii)). These results suggested that the key ACD regulators may not be required for erythroid enucleation *in vivo*. To formally exclude the possibility that a redundant physiological mechanism might act to rescue ACD defects *in vivo*, we investigated the ability of Par3, Scribble or Gpsm2-depleted orthochromatic erythroblasts to enucleate in media *ex vivo*[26] in the absence of surrounding supporting microenvironment. We found no differences in enucleation efficiencies of Par3, Scribble or Gpsm2 depleted erythroblasts



6	EpoR-Cre ⁺ ; Pard3 ^{+/+}	EpoR-Cre ⁺ ; Pard3 ^{fl/fl}	EpoR-Cre ⁺ ; Scrib ^{+/+}	EpoR-Cre ⁺ ; Scrib ^{f/fl}	Gpsm2 ^{+/+}	Gpsm2 ^{∆C/∆0}
Red blood cell [x10 ¹² /L]	4.59 ± 0.63	4.16 ± 0.97	4.84 ± 0.31	4.67 ± 0.24	5.07 ± 0.25	5.91 ± 0.10**
Hemoglobin [g/dL]	7.80 ± 0.59	6.58 ± 1.43	7.63 ± 0.48	7.42 ± 0.46	8.34 ± 0.29	7.60 ± 1.03
Hematocrit [L/L]	0.27 ± 0.03	0.23 ± 0.06	0.26 ± 0.03	0.26 ± 0.03	0.29 ± 0.01	0.29 ± 0.02
MCV [x10 ⁻¹⁵ /L]	58.73 ± 2.05	57.98 ± 1.87	54.07 ± 4.96	54.87 ± 4.86	56.98 ± 4.40	49.70 ± 3.27
RDW [%CV]	43.23 ± 3.19	45.78 ± 4.60	42.53 ± 0.68	44.07 ± 2.20	39.95 ± 3.08	38.47 ± 3.07



Fig 3. The ACD regulators Par3, Scribble and Gpsm2 are not required for erythroid enucleation during stress erythropoiesis. (A) Representative images of blood smears from peripheral blood derived from PHZ treated, agematched mice of the indicated genotypes of the different mouse models. Scale bar = 10µm. (B) Whole blood analysis using the CELL-DYN Sapphire System was performed on peripheral blood derived from PHZ treated, agematched mice of the indicated genotypes of the indicated mouse models. Data represent the mean (+/- SD) of 3–6 independent

experiments. (C) Orthochromatic erythroblasts were enriched by FACS (Aria II) from spleen of PHZ treated mice of the indicated genotypes of the indicated mouse models and incubated in 96-well plates at 30000 cells per well. Graphs showing enucleation efficiencies 5h post sort quantified by LSR II. Data represent the mean (+/- SD) of 3 independent experiments. *P< 0.05, **P< 0.01, ***P< 0.001, ***P< 0.001 (unpaired student's t-test). (D) Whole blood derived from PHZ treated, aged-matched mice of the indicated genotypes of the different mouse models was exposed to decreasing concentrations of sodium chloride (NaCl) and the degree of hemolysis measured colorimetrically. Data represent the mean (+/- SD) of 3 independent experiments.

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compared to age-matched wild-type counterparts isolated from bone marrow (Fig 1C), indicating that the key ACD regulators Par3, Gpsm2 or Scribble are not essential for enucleation.

Osmotic fragility is affected in erythrocytes with abnormal membrane integrity and/or abnormal surface area-to-volume ratios[30]. As many membrane proteins are asymmetrically distributed during the enucleation event, we also investigated the membrane integrity of erythrocytes depleted of Par3, Scribble or Gpsm2 by measuring osmotic fragility. However, no difference in the degree of hemolysis could be identified (Fig 1D (i-iii)).

To exclude the possibility that long-term physiological compensation in erythropoiesis was not masking enucleation defects, we investigated whether erythropoiesis in general is affected in animals harboring erythroblasts that lack ACD regulators. Analysis of bone marrow cellularity (Fig 2A, left) and the proportion of erythroid cells (Fig 2A, right) in femoral bone marrow, revealed no significant differences between mice deficient of erythroid Par3, Scribble or Gpsm2 expression compared to age-matched wild-type controls. Analysis of erythropoiesis during homeostasis in bone marrow also showed no significant difference in the percentage of the different developmental erythroblast stages (Fig 2B and 2C). Altogether, these data indicate that the Par3, Scribble and Gpsm2 polarity regulators are dispensable for the full maturation of erythrocytes *in vivo*.

Polarity regulators Par3, scribble and Gpsm2 are not essential for red blood cell development during stress erythropoiesis

As we did not observe any enucleation deficits in erythroblasts depleted of functional Par3, Scribble or Gpsm2 during steady-state conditions, we next evaluated whether loss of these ACD regulators impacted on enucleation during stress erythropoiesis as this could potentially reveal rate-limiting requirements for these genes in enucleation. However, no enucleation defects were observed in the different mouse models based on blood smear analysis (Fig 3A (i-iii)) and whole blood analysis (Fig 3B (i-iii)) following phenylhydrazine (PHZ) -induced hemolytic stress erythropoiesis. We did observe a small but significant increase in the number of erythrocytes in the peripheral blood of $Gpsm2^{\Delta C/\Delta C}$ mice, compared to age-matched controls but no other differences were observed (Fig 3A).

To test whether the *ex vivo* enucleating efficiency of orthochromatic erythroblasts isolated from spleens during stress erythropoiesis was affected by the functional loss of Par3, Scribble or Gpsm2, we compared enucleation rates of splenic orthochromatic erythroblasts from PHZ treated mice of the indicated genotypes to those of their age-matched controls (Fig 3C (i-iii)). Interestingly, the basal enucleation rate is much higher in splenic orthochromatic erythroblasts following PHZ induced stress erythropoiesis (around 60%) compared to the enucleation rate of orthochromatic erythroblasts isolated from bone marrow (up to 30%) (compare Fig 1C to Fig 3C). Regardless, no differences in enucleation efficiencies were observed between genotypes, suggesting that Par3, Scribble or Gpsm2 are not required for enucleation in the spleen during stress erythropoiesis. In addition, peripheral blood from PHZ treated mice lacking erythroid expression of Par3 or Scribble or whole body Gpsm2 were subjected to decreasing



Fig 4. The ACD regulators Par3, Scribble and Gpsm2 are not required for stress erythropoiesis in the spleen. (A) Graphs showing spleen weights relative to body weights determined from untreated and PHZ treated, age-matched mice of the indicated genotypes of the different mouse

models. Data represent the means (+/- SD) of 3–5 independent experiments. (B) Graphs showing total cell numbers per spleen (on left) and percentages of erythroid cells (Ter119 positive) in spleen (on right) isolated from PHZ treated, age-matched mice of the indicated genotypes of the different mouse models. Data represent the means (+/- SD) of 3–6 independent experiments. (C) Bar graphs showing percentages of erythroid cells at the different developmental stages (I-V) during stress erythropoiesis in the spleen from PHZ treated mice of the indicated genotypes of the different mouse models. Data represent the means (+/- SD) of 3–5 independent experiments. *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001 (unpaired student's t-test).

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concentrations of NaCl and the degree of hemolysis measured and compared to age-matched controls. Again, no difference in the degree of hemolysis could be identified (Fig 3D).

Analysis of stress erythropoiesis in general also showed no defects: Spleen weights revealed no significant differences in mice deficient of erythroid Par3, Scribble or Gpsm2 expression compared to age-matched wild-type controls (Fig 4A). Analysis of cellularity (Fig 4B, left) and the proportion of erythroid cells (Fig 4A, right) or their erythroid subsets (Fig 4C) in suspensions made from spleens isolated from PHZ treated mice, also revealed no significant differences. Collectively, these results suggest that enucleation is not affected by erythoid specific loss of functional Par3 or Scribble, or whole body loss of Gpsm2 under steady-state or stress conditions.

It is important to note that efficient erythropoiesis is crucial for organismal survival and that knockdown of functionally important genes is often associated with either embryonic death or high redundancy. We have started to examine compensatory mechanisms by qPCR (S3 Fig) but could not detect any significant compensatory upregulation in the knockout models tested here at a transcriptional level. However, it will be crucial to test whether other ACD regulators can compensate for the loss of Par3, Scribble or Gpsm2 in erythroid cells. The generation and analysis of double knockout mutants may shed light on this possibility. Future experiments will also involve examining expression and localization of ACD regulators at the protein level. Nevertheless, given that Par3, Scribble and Gpsm2 are fundamental regulators of polarity and ACD in invertebrates and mammalian cells, our results strongly indicate that the extreme asymmetry during erythroid enucleation is not established through these key ACD regulators. Red blood cells develop in erythroblastic islands, where erythroblasts differentiate bound to a central macrophage[31]. It is therefore possible that adhesion to the extracellular environment, e.g. Emp (erythroblasts macrophage protein) dependent adhesion between erythroblasts and macrophages [32, 33], or adhesion between erythroblasts and fibronectin [34], or possibly adhesion amongst erythroblasts[35], could provide the necessary cues to control asymmetric positioning of the cytoskeleton prior and during enucleation, rather than cellintrinsic asymmetry. The observation that erythroid enucleation can occur in vitro in the absence of many of these microenvironmental cues though also suggests that other non-deterministic and stochastic mechanisms could be utilized to generate these asymmetries. Indeed, a key difference between ACD and erythroid enculeation is that canonical ACD involves not just polarization, but also alignment of that polarity with the spindle. Further, instead of a specific directionality, the recruitment of the nucleus to any part of the erythrocyte cortex might be the only requirement. Together, our findings suggest that erythrocyte enucleation may only have a minimal requirement for extracellular cues and the coordination provided by polarity complexes.

Supporting Information

S1 Fig. ACD genes are expressed in orthochromatic erythroblasts. Gene expression of genes from Scribble (A), Gpsm2 (B) and Par (C) complex in orthochromatic compared to late, but still proliferating erythroblasts isolated from spleens of phenylhydrazine (PHZ) treated wild-type mice. Data are means (+/- SD) of 3 independent experiments. *P < 0.05, **P < 0.01,

P< 0.001, *P< 0.0001 (unpaired student's t-test). (TIF)

S2 Fig. Molecular characterization of mouse models. (A) To investigate the biological functions of polarity regulator Par3 in vivo, mice harboring a Pard3 construct flanked by loxP sites were crossed to $EpoR-Cre^+$ mice. (i.) Representative PCR analysis showing deletion of the floxed Pard3 allele in erythroblasts isolated by FACS (Aria II) from bone marrow (BM) suspensions harvested from mice of the indicated genotypes (ii.) Graph showing relative Pard3 gene expression measured in FACS sorted erythroid cells derived from bone marrow harvested from mice of the indicated genotypes. Data represent the mean (+/- SD) of 3 independent experiments. (B) Mice harboring a *Scrib* construct flanked by loxP sites were crossed to *EpoR*- Cre^+ mice. (i.) Representative PCR analysis showing deletion of the floxed Scrib allele in erythroblasts isolated by FACS (Aria II) from bone marrow (BM) suspensions harvested from mice of the indicated genotypes (ii.) Graph showing relative Scrib gene expression measured in FACS sorted erythroid cells derived from bone marrow harvested from mice of the indicated genotypes. Data represent the mean (+/- SD) of 2-3 independent experiments. (C) Characterization of Gpsm2^{ΔC} mice. (i.) Representative PCR analysis showing deletion of the *Gpsm2* allele in erythroblasts isolated by FACS (Aria II) from bone marrow (BM) suspensions harvested from mice of the indicated genotypes (ii.) Graph showing relative Gpsm2 gene expression measured in FACS sorted erythroid cells derived from bone marrow harvested from mice of the indicated genotypes. Data represent the mean (+/- SD) of 3 independent experiments. *P< 0.05, **P< 0.01, ***P< 0.001, ****P< 0.0001 (unpaired student's t-test). (TIF)

S3 Fig. Characterization of potential compensatory mechanisms. Gene expression of genes from Scribble, Gpsm2 and Par complex in erythroblasts isolated by FACS (Aria II) from bone marrow suspensions harvested from (**A**) *EpoR-Cre⁺;Pard3*^{fl/fl}, (**B**) *EpoR-Cre⁺;Scrib*^{fl/fl} and (**C**) *Gpsm2*^{AC} mice and their age-matched controls. Data are means (+/- SD) of 3 independent experiments. *P< 0.05, **P< 0.01, ****P< 0.001, ****P< 0.001 (unpaired student's t-test). (TIF)

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