

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27

**MRI Imaging of the Hemodynamic Vasculature of Neuroblastoma Predicts
Response to Anti-angiogenic Treatment**

Konstantinos Zormpas-Petridis¹, Neil P. Jerome^{1,2,3}, Matthew D. Blackledge¹, Fernando Carceller⁴, Evon Poon⁴, Matthew Clarke⁵, Ciara M. McErlean¹, Giuseppe Barone⁶, Alexander Koers⁴, Sucheta J. Vaidya⁴, Lynley V. Marshall⁴, Andrew D.J. Pearson⁴, Lucas Moreno⁷, John Anderson^{6,9}, Neil Sebire^{8,9}, Kieran McHugh¹⁰, Dow-Mu Koh¹, Yinyin Yuan⁵, Louis Chesler⁴, Simon P. Robinson¹ and Yann Jamin^{1,*}.

¹ Division of Radiotherapy and Imaging, The Institute of Cancer Research, London and The Royal Marsden NHS Trust, 15 Cotswold Rd. Belmont, Sutton, Surrey SM2 5NG, United Kingdom.

² Department of Circulation and Medical Imaging, Norwegian University of Science and Technology (NTNU), Trondheim, Norway

³ Clinic of Radiology and Nuclear Medicine, St. Olavs Hospital, Trondheim, Norway

⁴ Division of Clinical Studies, The Institute of Cancer Research, London and The Royal Marsden NHS Trust, 15 Cotswold Rd. Belmont, Sutton, Surrey SM2 5NG, United Kingdom.

⁵ Division of Molecular Pathology, The Institute of Cancer Research, London and The Royal Marsden NHS Trust, 15 Cotswold Rd. Belmont, Sutton, Surrey SM2 5NG, United Kingdom.

⁶ Department of Pediatric Oncology, Great Ormond Street Hospital for Children, London WC1N 3JH, United Kingdom

⁷ Clinical Research Unit, Pediatric Oncology, Hematology and Stem Cell Transplant Department, Hospital Infantil Universitario Niño Jesús, Madrid, Spain.

⁸ Department of Histopathology, Great Ormond Street Hospital for Children, London WC1N 3JH, United Kingdom

1 ⁹ UCL Great Ormond Street Institute of Child Health, London WC1N 1EH, UK

2 ¹⁰ Department of Radiology, Great Ormond Street Hospital for Children, London WC1N 3JH, United
3 Kingdom

4 **Corresponding author:**

5 Dr Yann Jamin, Centre for Cancer Imaging, Division of Radiotherapy and Imaging, The
6 Institute of Cancer Research, London, 15 Cotswold Rd. Belmont, Sutton, Surrey SM2 5NG,
7 UK. Email: yann.jamin@icr.ac.uk

8

9 **Research Article**

10

11 **Running title:** MRI predicts response to VEGFR-targeted therapy in neuroblastoma.

12

13 **Keywords:** MRI, Neuroblastoma, MYCN, VEGF, Biomarkers

14

15 **Financial support:** Cancer Research UK and EPSRC to the Cancer Imaging Centre at ICR,
16 in association with the MRC and Department of Health (England) (C1060/A10334 and
17 C1060/A16464), NHS funding to the NIHR Biomedicine Research Centre and the Clinical
18 Research Facility in Imaging, the NIHR GOSH Biomedical Research Centre, The Rosetrees
19 Trust (KZP), Children with Cancer UK (E.P.). Cancer Research UK Programme Grant
20 (C18339) (L.C.). L.M. was funded by the Oak Foundation to the Royal Marsden and by
21 Instituto de Salud Carlos III through Rio Hortega CM12/00260 and Juan Rodes JR15/00041
22 contracts. J.A. is supported by the NIHR GOSH Biomedical Research Centre and GOSHCC
23 research leadership award. YJ is a Children with Cancer UK Research Fellow.

24

25 **Conflict of interest:** The authors have no conflict of interest to disclose

26

27

28 **Word count:** 4990

29

30 **Number of tables:** 0

31

32 **Number of figures:** 7

33

1 **Abstract**

2

3

4 Childhood neuroblastoma is a hypervascular tumor of neural origin for which antiangiogenic
5 drugs are currently being evaluated; however, predictive biomarkers of treatment response,
6 crucial for successful delivery of precision therapeutics, are lacking. We describe a MRI-
7 pathological cross-correlative approach using intrinsic susceptibility (IS-) and susceptibility
8 contrast (SC-) MRI to non-invasively map the vascular phenotype in neuroblastoma Th-
9 MYCN transgenic mice treated with the vascular endothelial growth factor receptor inhibitor
10 cediranib. We showed that the transverse MRI relaxation rate R_2^* (s^{-1}) and fractional blood
11 volume fBV (%) were sensitive imaging biomarkers of hemorrhage and vascular density,
12 respectively, and were also predictive biomarkers of response to cediranib. Comparison with
13 MRI and pathology from patients with MYCN-amplified neuroblastoma confirmed the high
14 degree to which the Th-MYCN model vascular phenotype recapitulated that of the clinical
15 phenotype, thereby supporting further evaluation of IS- and SC-MRI in the clinic. This study
16 reinforces the potential role of functional MRI in delivering precision medicine to children
17 with neuroblastoma.

18

19

20 **Significance**

21

22 This study shows that functional MRI predicts response to vascular-targeted therapy in a
23 genetically-engineered murine model of neuroblastoma.

1 **Introduction**

2

3 Neuroblastoma is a tumor of childhood that arises within the embryonic sympatho-adrenal
4 lineage of the neural crest. Neuroblastoma in high-risk form is chemoresistant, and metastatic
5 relapse still accounts for 13% of all cancer-related death in children. Amplification of the
6 proto-oncogene *MYCN* is the commonest genomic aberrations and defines a subgroup of
7 children at high-risk of developing uncontrollable refractory or relapsing disease despite
8 aggressive multimodal therapy (1).

9

10 In neuroblastoma, a high vascular index is associated with increased disseminated disease,
11 amplification of *MYCN*, unfavorable histology and overall poor prognosis (2). Increased
12 microvascular proliferation and other specific vascular morphological patterns are associated
13 with even poorer prognosis, highlighting the pivotal role of angiogenesis in determining the
14 clinical behavior of neuroblastoma (3,4). The vascular endothelial growth factor (VEGF)
15 family is a key regulator of angiogenesis in neuroblastoma and high VEGF expression at the
16 time of diagnosis is associated with poor outcome (5). Numerous anti-angiogenic therapies
17 are being evaluated in early phase pediatric clinical trials in children with solid tumors,
18 including the anti-VEGF monoclonal antibody bevacizumab in combination with
19 temozolomide (BEACON, NCT02308527), and the tyrosine kinase inhibitors regorafenib
20 (NCT02085148) and pazopanib as a single agent (NCT01956669) or in combination with
21 metronomic oral topotecan (TOPAZ), as well as the panVEGFR inhibitor cediranib currently
22 being evaluated in children with metastatic alveolar soft part sarcoma (NCT00942877).

23

24 Magnetic resonance imaging (MRI) is becoming the preferred clinical imaging technique for
25 the management of children with neuroblastoma because of its exquisite soft tissue contrast.
26 MRI provides excellent anatomical information at diagnosis and follow up while sparing
27 exposure to ionising radiation associated with computed tomography (CT) (6). Advanced
28 functional MRI techniques can additionally be used to define quantitative imaging
29 biomarkers that inform on biologically relevant structure-function relationships in tumors *in*
30 *vivo*. Perfusion MRI (pMRI) methodologies such as dynamic contrast enhanced (DCE-) MRI
31 using low molecular weight gadolinium chelates are often used to evaluate vascular response
32 to VEGF signaling inhibitors in adult oncology clinical trials. However, these techniques
33 suffer from marked measurement variability and are challenging to perform in young
34 children (7,8). Arterial spin labeling (ASL-) MRI is an emerging and attractive contrast-free

1 approach for the evaluation of cerebral perfusion, yet many challenges need to be overcome
2 before its utility to assess vascular perfusion in extra-cranial tumors can be evaluated. Two
3 complementary magnetic susceptibility-based MRI approaches are being actively exploited to
4 assess tumor vascular function and response. Tumor vasculature has been studied using *i*)
5 intrinsic susceptibility (IS-) MRI, which measures the intrinsic transverse relaxation (R_2^*)
6 contrast produced by paramagnetic deoxyhemoglobin within tumor blood vessels, and *ii*)
7 susceptibility contrast (SC-) MRI, which relies on the intravenous administration of
8 ultrasmall superparamagnetic iron oxide (USPIO) particles. Distribution of USPIO particles
9 within tumors causes regional increases in tumor R_2^* (ΔR_2^*), from which the fractional blood
10 volume (fBV , %) can be derived (9,10). The long intravascular half-life of USPIO particles
11 enables steady state acquisition and high-resolution mapping of regional variations in tumor
12 perfusion. Both R_2^* and fBV has been shown to be sensitive imaging biomarkers of response
13 to vascular-targeted therapies *in vivo* in the preclinical setting (11-13).

14

15 Imaging biomarkers must undergo stringent validation before they can be deployed clinically
16 (14). Early imaging biomarker development demands close imaging-pathology correlation, to
17 understand the biological processes underpinning the imaging measurement, which can be
18 meaningfully studied using animal models. The Th-*MYCN* genetically-engineered mouse
19 (GEM) model of neuroblastoma has been shown to faithfully recapitulate the major genetic
20 and patho-physiological features of the childhood disease (12,15). The primary abdominal
21 tumors in the Th-*MYCN* mice present with multiple para- and intra- tumoral anatomical
22 landmarks detectable by conventional MRI, which can be used to ensure the accurate
23 registration between the functional MRI-derived parametric maps and histopathology crucial
24 for the validation of imaging biomarkers. The GEM models of neuroblastoma thus represent
25 an information-rich platform with which to evaluate promising therapeutic strategies and
26 associated non-invasive imaging biomarkers.

27

28 In this study, we investigated the utility of SC- and IS-MRI to inform on the functional tumor
29 vasculature, and its response to the potent pan-VEGFR inhibitor cediranib, in tumors arising
30 in the Th-*MYCN* GEM model of neuroblastoma. We then compared the regional distribution
31 of tumor fBV and R_2^* with aligned whole-slide digitized pathology stained for vessel density
32 and red blood cell (RBC) aggregation respectively, and validated, via hotspot mapping and
33 spatial statistics, to determine whether the MRI biomarkers can robustly characterize the

1 vascular phenotype of neuroblastoma *in vivo*. We also demonstrate here that both baseline
2 fBV and R_2^* are predictive imaging biomarkers of tumor response to cediranib. The
3 pathological attributes associated with the differential sensitivity to cediranib treatment in the
4 Th-*MYCN* model are discussed and positioned against available clinical MRI findings of
5 *MYCN*-amplified neuroblastoma, including our initial experience with IS-MRI.
6

1

2 **Materials and Methods**

3

4 ***Animals***

5 All experiments were approved by The Institute of Cancer Research Animal Welfare and
6 Ethical Review Body and performed in accordance with the UK Home Office Animals
7 (Scientific Procedures) Act 1986, the United Kingdom National Cancer Research Institute
8 guidelines for the welfare of animals in cancer research (16) and the ARRIVE (animal
9 research: reporting in vivo experiments) guidelines (17).

10

11 Transgenic Th-*MYCN* mice were genotyped to detect the presence of the human *MYCN*
12 transgene (18). The study was performed using both male and female hemizygous mice,
13 which developed palpable tumors at 50–130 days with a 25% penetrance. Tumor
14 development was monitored weekly by palpation by an experienced animal technician. Mice
15 with palpable tumors were then enrolled (Day 0) and their tumor volume was subsequently
16 monitored by MRI. A total of 68 mice were enrolled with a median tumor volume of 801 ± 63
17 mm^3 (median ± 1 s.e.m., ranging from 143 to 2055 mm^3). Mice were housed in specific
18 pathogen-free rooms in autoclaved, aseptic microisolator cages (maximum of 4 mice per
19 cage).

20

21 ***Preclinical study design***

22 *Study 1.* We first evaluated the sensitivity of MRI to cediranib-induced acute modulation of
23 neuroblastoma vasculature. IS- and SC-MRI were performed prior to (*Day 0*) and 24 hours
24 after treatment started (*Day 2*). Mice were treated on *Day 1* with 6 mg/kg of cediranib p.o.
25 (obtained under material transfer agreement with AstraZeneca, n=10) or vehicle (n=8).
26 One mouse was excluded due to failed remote contrast injection. The 24h imaging
27 timepoint was chosen based on *i*) preliminary evidence that cediranib does not elicit any
28 significant volume reduction at this timepoint in the highly chemosensitive Th-*MYCN*
29 model (12) and *ii*) cediranib caused significant reductions in DCE-MRI parameters at this
30 timepoint in the adult Phase I clinical trial (19).

1 *Study 2.* In an additional cohort (n=12) we further evaluated the effect of sustained daily
2 treatment, with IS- and SC-MRI performed prior to, 24 hours and 7 days after daily
3 treatment with 6 mg/kg cediranib. MRI data were not collected from 2 mice at the 24h
4 timepoint.

5 *Study 3.* Guided by the results of Study 2, we subsequently acquired IS-MRI data from
6 additional Th-*MYCN* mice (n=25) prior to daily treatment with cediranib for 7 days
7 (bringing the total number of mice from which pre-treatment R_2^* data was acquired to 37).
8 The volumetric response to cediranib over 7 days treatment was monitored by T_2 -weighted
9 MRI only, and compared with that from mice treated daily with vehicle (n=12).

10

11 ***MRI***

12 All MRI studies were performed on a 7T Bruker horizontal bore MicroImaging system
13 (Bruker Instruments, Ettlingen, Germany) using a 3cm birdcage volume coil. Anesthesia was
14 induced by an intraperitoneal 5ml/kg injection of a combination of fentanyl citrate
15 (0.315mg/ml) plus fluanisone (10mg/ml) (Hypnorm, Janssen Pharmaceutical, Oxford, UK)
16 and midazolam (5mg/ml) (Roche, Welwyn Garden City, UK) and water (1:1:2). A lateral tail
17 vein was cannulated with a 27G butterfly catheter (Hospira) for remote administration of
18 USPIO particles. Core temperature was maintained at $\sim 37^\circ\text{C}$ with warm air blown through
19 the magnet bore.

20

21 For all the mice, anatomical T_2 -weighted transverse images were acquired from twenty
22 contiguous 1 mm-thick slices through the mouse abdomen, using a rapid acquisition with
23 refocused echoes (RARE) sequence with 4 averages of 128 phase encoding steps over a 3×3
24 cm field of view, an echo time (TE) of 36 ms, a repetition time (TR) of 4.5 s and a RARE
25 factor of 8. These images were used to determine tumor volumes, and for planning the
26 subsequent functional MRI measurements, which included optimization of the local field
27 homogeneity using FASTmap algorithm and the measurement of the baseline transverse
28 relaxation rate R_2^* (s^{-1}), which is sensitive to the concentration of paramagnetic species,
29 principally deoxyhemoglobin. R_2^* was quantified using a multiple gradient-recall echo
30 (MGE) sequence with 8 averages, and an acquisition time of 3 min 20 s. Images were
31 acquired using 8 echoes spaced 3 ms apart, an initial TE of 6 ms, a flip angle $\alpha=45^\circ$ and a TR
32 of 200 ms. A dose of 150 $\mu\text{mol Fe/kg}$ of the USPIO particle preparation P904 (overall

1 particle size ~25–30 nm diameter, Guerbet) was then administered intravenously. After 3
2 minutes to allow for equilibration, a second set of identical MGRE images acquired.

3
4 All the MRI data were acquired with a matrix size of 128x128 over a 3×3 cm field of view.
5 Tumor volumes were determined using segmentation from regions of interest drawn on T₂-
6 weighted images for each tumor-containing slice using OsiriX. Tumor R₂* maps were
7 calculated from regions of interest drawn for each tumor-containing slice from the MGRE
8 images acquired prior to and following administration of USPIO particles by fitting a single
9 exponential to the signal intensity echo time curve on a voxel-by-voxel basis using a robust
10 Bayesian approach using in-house software (ImageView, developed in IDL, ITT Visual
11 Information Systems, Boulder, Colorado, USA). Parametric maps of tumor fBV (%) were
12 subsequently calculated using the USPIO-induced change in R₂* (ΔR_2^*), as described
13 previously (20).

15 ***Tissue preparation for histology***

16 Guided by T₂-weighted anatomical images, tumors were carefully excised and orientated for
17 subsequent histopathological processing. Formalin-fixed and paraffin embedded tumors were
18 cut in 3 μ m sections and were stained with hematoxylin and eosin (H&E) and for the murine
19 vascular endothelial marker endomucin (rabbit EP3095, Millipore, Watford, UK). Whole-
20 Slide images were digitized using a Hamamatsu NanoZoomer XR scanner (20x
21 magnification, 0.46 μ m resolution, Hamamatsu, Japan).

23 ***MRI-histology alignment***

24 The MRI slice of interest were visually aligned with digitized whole-slide H&E stained
25 images using anatomical landmarks from T₂-weighted images, including, the shape of the
26 tumor, the position of the kidneys, the position and/or orientation of tumor-displaced and –
27 circumferentially surrounded abdominal aorta and renal aorta/vein and the presence of lymph
28 nodes.

29

30

1 ***Computational pathology***

2 Both red blood cells (RBC) and endomucin staining were automatically extracted from T₂-
3 weighted MRI corresponding digitized whole-slide histopathological images for the
4 validation of R₂* and fBV as biomarker of red blood cells aggregation and vascular
5 perfusion. Histological images (0.46x0.46μm pixel resolution) were first split into tiles of
6 2000x2000 pixels (jpeg) for computational efficiency.

7

8 *RBC extraction from HE-stained sections.* A macro was written in Fiji to extract RBC from
9 each tile by applying color deconvolution to extract the eosin color channel followed by the
10 application of Otsu's automatic threshold detection method, both using ImageJ/Fiji plugins
11 (with Java 8) (21). Algorithm accuracy was tested using independent annotation of 561 RBC
12 and 591 non-RBC points in 9 samples.

13

14 *Endomucin extraction from endomucin-stained IHC sections.* A macro was written in Fiji to
15 extract endomucin from each tile by applying color deconvolution to extract the brown color
16 channel followed by the application of Maximum Entropy threshold detection method, both
17 using ImageJ/Fiji plugins (Java 8). Algorithm accuracy was tested using independent
18 annotation of stained/non-stained points (688/734) on 9 different samples.

19

20 *Generation of RBC and endomucin parametric maps.* Whole-slide images of RBC and
21 endomucin staining were converted into binary and processed to match MRI resolution
22 (234x234μm), with the fraction of pixels occupied by RBC or endomucin staining within
23 518x518 pixel-regions representing a single pixel in the final RBC map.

24

25 *Comparison of R₂* vs RBC and fBV vs endomucin parametric maps* was performed visually
26 following the application the kernel density estimation (KDE) to display hotspots of high,
27 above the 85th percentile for each tumor sample, R₂* and RBC (%), fBV (%) and endomucin
28 staining (%) values using the 'MASS' package in R (22,23).

29

30

1 ***MRI-histology registration***

2 *Coherent point drift (CPD) registration algorithm.* Accurate registration between histology
3 and MRI images is essential to derive robust conclusions and apply spatial statistics. We
4 applied the automatic CPD algorithm for both *i)* rigid and *ii)* non-rigid registration to
5 corresponding pairs of images (R_2^* or fBV maps and computed maps of red blood cells
6 (RBC) and endomucin respectively) in Matlab (24). This algorithm was primarily selected
7 because it preserves topological structures due to the coherent motion of the point sets. To
8 avoid any bias, the point sets (features) from each map were edges extracted using a simple
9 Canny edge detector. Independently, we also performed *iii)* a manual registration by rotation
10 and scaling to match.

11

12 ***Statistical analysis***

13 Statistical analysis was performed with GraphPad Prism 6 (GraphPad Software Inc., La Jolla,
14 USA). The mean of median values for all the quantitative MRI parameters, the mean values
15 for tumor volume, RBC and endomucin were used for statistical analysis with a 5% level of
16 significance. Any significant differences in tumor volume and quantitative histopathological
17 parameters were identified using Student's 2-tailed unpaired t-test, with a 5% level of
18 significance. Significant correlations between the mean values. All linear correlations were
19 determined by Pearson's correlation method. Level of significance was 5%.

20

21 *Spatial statistics.* The Mantel test, a non-parametric analysis of associations between
22 corresponding positions of two distance matrices, was performed on registered pairs of MRI
23 maps and computed maps of histopathological features using the "ade4" library in R. Distance
24 matrices were calculated using the "dist" function.

25

26 ***Clinical investigations***

27 Based on the results of the preclinical investigations we aimed to *i)* evaluate the clinical
28 relevance of the vascular phenotype observed in the Th-MYCN tumors by reviewing
29 available diagnostic MRI data from patients with *MYCN*-amplified neuroblastoma, and *ii)*
30 evaluate the feasibility of acquiring IS-MRI in children with neuroblastoma.

31

1 Informed consent was obtained from all parents or patients following IRB approval of the
2 prospective study with IS-MRI at the Royal Marsden Hospital. For the retrospective review
3 of routine diagnostic imaging at Great Ormond Street Hospital, anonymised data was
4 provided to the researchers.

5

6 *Retrospective analysis of MRI scans performed at initial diagnosis.* Patients (n=19, 13 male/
7 6 female) with confirmed *MYCN*-amplified disease included in this retrospective analysis
8 were 1 day to 3.6 years old and imaged between May 2008 and July 2017 at Great Ormond
9 Street Hospital. Exclusion criteria were *i*) no MRI at diagnosis or *ii*) MRI performed
10 externally. Diagnostic imaging was performed on a 1.5 T Magnetom Avanto system
11 (Siemens Healthcare, Erlangen, Germany) and included short time inversion recovery (STIR)
12 T₂-weighted sequences (TE=60ms, TR=8000ms) and spectral attenuated inversion recovery
13 (SPAIR) T₁-weighted BLADE sequence (TE=20ms, TR=600ms), prior and after the injection
14 of Gd chelate based contrast agent, diffusion-weighted MRI sequence (TE=90ms,
15 TR=2700ms, 5 b-values, b=0, 50-800s/mm²). Slice thickness was 6mm and the field of view
16 was adapted to each patient.

17

18 *Prospective study of the feasibility of IS-MRI in patients with refractory/relapsing*
19 *neuroblastoma.* The 3 patients (n= 2 male/ 1 female) with refractory/relapsing disease
20 included in this study were 5 to 7 years old and scanned between July 2013 and March
21 2015. MRI was performed on a 1.5 T Magnetom Avanto system using a phased-array body
22 coil and included T₂-weighted sequence (turbo spin echo sequence TE=80ms, TR=7000ms)
23 and T₁-weighted (TE=3ms, TR=10000ms) prior and after (last scan of the session) the
24 injection of Gd chelate based contrast agent, and a Diffusion-weighted MRI sequence
25 (TE=70ms, TR=3400ms, 6 b-values, b=0, 50-800s/mm²). IS-MRI was performed in 2.5 min
26 using a 2D multiple gradient-recall echo sequence (initial TE 5ms, 5 echoes, 10 ms apart,
27 TR= 102 ms, flip angle= 40°). Images were analysed and R₂* quantified in Matlab.

28

1 **Results**

2

3 *Baseline tumor fBV predicts response to cediranib in the Th-MYCN GEM model of*
4 *neuroblastoma.*

5 We first assessed changes in vascular function in Th-MYCN mice following treatment with
6 cediranib. At the time of enrolment, the mean tumor fBV quantified using SC-MRI in the Th-
7 MYCN mice (n=19) was $19.6 \pm 1\%$, consistent with the high vascular index of neuroblastoma.
8 Treatment with cediranib caused an acute and highly significant reduction at 24 hours in
9 tumor fBV ($\Delta fBV_{\text{cediranib}_{24h}} = -28 \pm 4\%$ vs $\Delta fBV_{\text{vehicle}_{24h}} = -6 \pm 8\%$, $p=0.0008$), which was
10 accompanied by a highly significant cytotoxic response ($\Delta \text{Volume}_{\text{cediranib}_{D0-D2}} = 18 \pm 8\%$ vs
11 $\Delta \text{Volume}_{\text{vehicle}_{D0-D2}} = 53 \pm 11\%$, $p=0.0006$, Fig 1. A-F). Sustained daily treatment over 7 days
12 caused a further significant reduction in fBV and anti-tumor activity (Fig 1. G-I). The average
13 baseline fBV strongly correlated with the cediranib-induced reduction in fBV ($r = -0.83$,
14 $p=0.0008$; Fig 1.J), and the value of fBV at baseline correlated with the subsequent cediranib-
15 induced tumor volumetric response at day 7 ($r = -0.65$, $p=0.02$; Fig 1.K).

16

17 *Baseline tumor R_2^* predicts response to cediranib in the Th-MYCN GEM model of*
18 *neuroblastoma.*

19 The analysis of the cohort of mice in which IS-MRI was routinely performed pre-treatment
20 confirmed that cediranib effectively suppresses the aggressive tumor growth that is typical of
21 this model ($\Delta \text{Volume}_{\text{cediranib}_{D0-D7}} = -16 \pm 5\%$ vs $\Delta \text{Volume}_{\text{vehicle}_{D0-D7}} = 142 \pm 11\%$, $p < 0.0001$,
22 $n=37$ and $n=12$). However, a range of volumetric response, from progressive disease to
23 partial response, was observed (Fig 2.A). Examination of our pre-treatment T₂-weighted
24 anatomical images and native R_2^* parametric maps revealed that responsive tumors exhibited
25 a characteristically heterogeneous appearance, with areas of hypointense T₂ signal and
26 relatively fast R_2^* ($R_2^*_{\text{mean}} = 113 \pm 4 \text{ s}^{-1}$) (Fig 2.B&C), whereas progressive tumors typically
27 demonstrated a more homogeneous, isointense appearance on T₂ and significantly slower
28 native R_2^* values ($R_2^*_{\text{mean}} = 68 \pm 5 \text{ s}^{-1}$, $p < 0.0001$). Cediranib-induced changes in tumor volume
29 over 7 days treatment correlated with native tumor R_2^* measured before treatment ($r = -0.72$,
30 $p < 0.0001$, Fig 2.D).

31

1 *Spatial heterogeneity in fBV reflects regional variations in the microvasculature of*
2 *neuroblastoma.*

3 The numerous anatomical landmarks (kidney, spleen, abdominal arteries, lymph nodes)
4 clearly evident on T₂-weighted MRI were used to guide the careful excision and subsequent
5 orientation of digitized pathology with SC- and IS-MRI-derived parametric maps. Intra-
6 tumoral regional differences in fBV visually reflected spatial variations in computed maps of
7 vascular endothelial endomucin staining, automatically extracted from high-resolution IHC
8 images with an accuracy of 97%, after automatic registration using the coherent point drift
9 (CPD) algorithm (Fig 3). This was corroborated by the location, size, shape and orientation of
10 hotspots identified on kernel density estimation (KDE) maps of high values (above the 85th
11 percentile of each tumor sample) of fBV and endomucin (Fig 4. A). Application of the
12 Mantel statistical test revealed significant ($p < 0.05$) correlation ($0.10 < r < 0.58$,
13 supplementary table 1) between the distance matrices of fBV and endomucin in 16 out of 23
14 tumors. Endomucin staining fraction corroborated the decrease in fBV measured
15 longitudinally over 7 days, with significantly lower tumor values determined in the cohort
16 treated with cediranib for 7 days than in the cohort of animals treated with a single dose of
17 cediranib; both cohorts demonstrated lower values of fBV compared with the vehicle control
18 group (Fig 4. B&C). Calculated tumor median values of fBV strongly correlated with median
19 values of endomucin fraction area across the cohorts ($r = 0.85$, $p < 0.0001$) (Fig 4.D).

20

21 *Regional heterogeneity in R₂* reflects variations in RBC distribution in neuroblastoma*

22 Regional differences in parametric R₂* maps visually reflected spatial variations in
23 extravasated RBC aggregation or RBC-filled necrotic areas (defined as large areas of cell
24 damage) in the digitized maps, which was corroborated by the location, size, shape and
25 orientation of hotspots on KDE maps of high values (above the 85th percentile of each tumor
26 sample) of R₂* and RBC, automatically extracted from high-resolution H&E-stained images
27 with an accuracy of 99.7%, after automatic registration using the CPD algorithm (Fig 5). The
28 Mantel statistical test revealed significant ($p < 0.05$) spatial correlation ($0.10 < r < 0.76$,
29 Supplementary table 2) between the distance matrices of R₂* and RBC in 9 out of 12 tumors.
30 Calculated tumor median values of R₂* strongly correlated with the mean values of RBC
31 fraction area ($r = 0.87$, $p < 0.0003$). Tumor median R₂* and mean value of RBC fraction also
32 correlated with endomucin staining ($r = 0.53$, $p = 0.08$ and $r = 0.64$, $p = 0.02$, respectively).

33

1

2 *Innate resistance to cediranib is associated with a differentiating phenotype*

3 We then performed a semi-quantitative analysis to calibrate the tumor R_2^* phenotype against
4 pathological evaluation of aligned H&E stained sections (Fig 6, Supplementary table 3). For
5 tumors with a median R_2^* between ~ 80 and $\sim 110s^{-1}$, increasing tumor median R_2^* reflected
6 increasing area/proportion of the tumor presenting with a hemorrhagic and hypervascular
7 phenotype, characterized by large sinusoidal-shaped vessels. In comparison, non-
8 hemorrhagic regions remained well-vascularized but with markedly reduced hemorrhage and
9 smaller capillary-like shape vessels. In these tumors, the extravasated RBCs conserved their
10 biconcave shape and appeared intact. Tumors with a median $R_2^* > 110s^{-1}$ additionally
11 presented with large necrotic regions filled with both intact and damaged RBCs. Tumors
12 exhibiting a median $R_2^* < 70s^{-1}$, (comparable to tumors which progressed while on
13 cediranib), presented a very different phenotype, characterized by the absence of hemorrhage
14 (including in necrotic areas), reduced microvessel density with smaller more regular
15 capillaries-like vessels, and large regions of differentiating neuroblasts (yet with only very
16 few mature ganglion cells) arranged in islands, separated by a large amount of neuropil.

17

18 *MYCN-amplified childhood neuroblastomas are hemorrhagic*

19 To understand the relevance of the tumor MRI phenotypes observed in the Th-*MYCN* GEM
20 mouse to *MYCN*-driven childhood neuroblastoma, we reviewed anatomical T_2 -weighted, T_1 -
21 weighted and contrast enhanced (CE-) T_1 -weighted images, as well as diffusion-weighted
22 (DW-) MRI-derived apparent diffusion coefficient (ADC) maps, acquired from 19 patients
23 with *MYCN*-amplified neuroblastoma at the time of diagnosis. All patients presented with
24 abdominal tumors showing a high degree of heterogeneity (both hyper- and hypo-intensity)
25 on T_2 -weighted MRI and regional hyper-intensity on T_1 -weighted MRI (Figure 7.A-E). Based
26 on the well-established appearance of ageing blood on both T_2 - and T_1 -weighted MRI in
27 hematoma (Figure 7.F), this appearance suggests the presence of blood products at different
28 stage of hemoglobin degradation. Most tumors presented a central region which did not
29 enhance on CE-MRI and showed elevated ADC values, suggesting the presence of large
30 areas of non-viable tissue. Additionally, a few tumors presented with cystic components,
31 with/without blood. Interestingly, 5 out of the 19 patients also presented with contiguous or
32 satellite masses showing homogeneous signal on both T_1 (isointense) and T_2 (hyperintense)
33 MRI, homogeneous contrast enhancement and low ADC values, indicative of water

1 restriction (Figure 7.B and C). This MRI phenotype suggests a non-hemorrhagic and cellular-
2 dense tumor phenotype within these lobes.

3 We also compared the radiology of two cases of ganglioneuroblastoma: a
4 ganglioneuroblastoma intermixed (GNB) and ganglioneuroblastoma nodular (GNBn), two
5 entities that are uncommonly MYCN-amplified. Yet GNB nodular are characterized by the
6 presence of macroscopic undifferentiated neuroblastoma nodule(s) of Schwannian stroma-
7 poor components coexisting with ganglioneuroblastoma-intermixed of stroma- rich
8 component or ganglioneuroma (fully differentiated) of stroma-dominant component. The
9 neuroblastoma nodule in the GNBn was easily identifiable with low ADC values, consistent
10 with a stroma-poor neuroblastoma phenotype, and demonstrated marked hypointensity on T₂-
11 weighted images, hyper-intensity on T₁-weighted images and high gadolinium avidity,
12 overall suggesting that the neuroblastic nodule presented a highly vascular and hemorrhagic
13 phenotype.

14 *Initial experience with IS-MRI in children with neuroblastoma*

15 Based on the initial results obtained in the Th-MYCN model, we have introduced IS-MRI into
16 clinical trials in patients with relapsed/refractory neuroblastoma as a proof of concept. Our
17 initial experience based on three patients scanned at 1.5T (Figure 7.G) showed that tumors
18 (n=5) presented with a heterogeneous regional distribution of R₂^{*}, with tumor median R₂^{*}
19 values ranging from 20 to 41s⁻¹ (Figure 7.H), a range similar to the one measured in mice
20 when corrected for the difference of magnetic field strength. Regionally, higher R₂^{*} values
21 were associated with hypo-intensity on T₂-weighted images, higher gadolinium avidity and
22 lower ADC values. Combined with the apparent hemorrhagic nature of MYCN-amplified
23 neuroblastoma, this pilot study suggests that IS-MRI should be prospectively evaluated in
24 children with neuroblastoma to determine its clinical utility as an imaging biomarker.

1 Discussion

2

3 There is a current paradigm shift toward a stratified precision medicine approach for children
4 with cancer. Central to this approach is the integration of both genomic and biological
5 information including pharmacodynamic and predictive biomarkers of response to
6 incorporate urgently-needed novel therapies into treatment schedules. With increasing
7 evidence for the contribution of angiogenesis in determining and predicting the biological
8 behavior of neuroblastoma, therapeutic approaches targeting the unique vascular phenotype
9 of neuroblastoma are being accelerated in early phase pediatric clinical trials. However, there
10 are currently no validated biomarkers predicting the clinical response or therapeutic benefits
11 to anti-angiogenic therapy.

12

13 Whilst MRI is becoming the preferred clinical imaging method in the management of
14 children with neuroblastoma particularly at diagnosis and for routine follow-up, the unique
15 ability of advanced functional MRI techniques to non-invasively quantify changes in tumor
16 vascular architecture and function has not yet been fully exploited (25). In this study, we have
17 utilized the multiple intra- and para- anatomical tumor landmarks detectable on conventional
18 T₂-weighted MRI in the Th-*MYCN* mouse, together with the CDP algorithm for registration
19 to develop a robust MRI-histopathology correlation pipeline. The use of KDE hotspot
20 mapping and non-parametric Mantel statistical test demonstrated that *f*BV, measured *in vivo*
21 by SC-MRI, spatially correlates with stained area for the endothelial cell marker endomucin
22 in tumors arising in the Th-*MYCN* mouse model of neuroblastoma. This approach thus
23 validates *f*BV as a sensitive and but also a specific imaging biomarker of tumor vascular
24 perfusion, and its therapeutic modulation by the potent VEGFR inhibitor cediranib.
25 Furthermore, we found a negative correlation between tumor *f*BV and cediranib-mediated
26 changes in *f*BV, suggesting baseline *f*BV as a potential predictive biomarker of vascular
27 response to VEGFR inhibition. Importantly, our data suggest that both baseline tumor *f*BV
28 and R₂* are predictive of the longer-term tumor response to VEGFR-targeted therapies such
29 as cediranib.

30 Several pre-clinical studies have demonstrated that quantitation of tumor *f*BV using SC-MRI
31 provides a sensitive biomarker of response to vascular-targeted agents *in vivo* (10,11,13). The
32 data herein highlights the potential of SC-MRI to assess neuroblastoma vascular response to
33 VEGF signaling inhibitors (26-29), and to MYCN-targeted therapeutics whose mechanism of

1 action is predicted to elicit an anti-angiogenic effect (30-33). For example, direct silencing of
2 *MYCN* through bromodomain extra terminal (BET) domain inhibition, destabilization of
3 *MYCN* protein via the selective targeting of Aurora A kinase or mTOR/PI3k signaling, and
4 inhibition of anaplastic lymphoma kinase (ALK) and RET kinases (reviewed in (34)) have all
5 been linked with angiogenic blockade in aligned preclinical trials against neuroblastoma or
6 other pediatric solid tumors (31,32,35-37).

7 The high vascular permeability and density present in the Th-*MYCN* tumors and detected by
8 IS- and SC-MRI illustrate the archetypal VEGF-driven vascular remodeling triggered by
9 members of the *MYC* family oncogenes (31,38). There is increasing evidence that enhanced
10 angiogenic signaling is part of a wider *MYC*-driven adaptive program to nutrient deprivation,
11 to which *MYC*-driven tumors rapidly develop an addiction to maintain their growth (38,39).
12 As a result, any blockade of this program via *MYC* deactivation, metabolic inhibition or
13 vascular blockade triggers a wave of apoptosis, resulting in the unique rapid tumor debulking
14 observed in *MYC*-driven tumors (38,39). The correlation between baseline fBV and R_2^* and
15 cediranib-induced tumor debulking established in this study illustrates a similar dependence
16 on *MYCN*-driven angiogenic and vascular remodeling in the Th-*MYCN* GEM model. In
17 contrast, tumors at the lower range of baseline R_2^* and fBV values, which progress despite
18 cediranib treatment, presented with a more normalized vasculature. We also identified tumors
19 with large areas of differentiating neuroblasts, a phenotype similar to that in *MYCN*-driven
20 GEM models of neuroblastoma harboring constitutive activation of *ALK*, and in which
21 reduced VEGFR signaling has been reported (30,40,41). Clinically, differentiated
22 neuroblastoma displays low vascularity (2,3), partly due to the release strong anti-angiogenic
23 factors which directly antagonize the effect of VEGF (42,43). Additionally, in differentiating,
24 schwannian-stroma poor neuroblastoma, VEGF-mediated cross talk between differentiating
25 neuroblasts and endothelial cells has been identified which mimics normal vascular
26 maturation in the developing nervous system (44). Overall in this study, reduced tumor
27 sensitivity to cediranib was associated with a more stable vascular phenotype, suggesting a
28 reduced dependence of these non-responsive tumors on VEGF signaling to sustain growth.

29 Genetically-engineered models of cancer have been shown to faithfully recapitulate the major
30 molecular and histopathological features of human cancer, and to have a strong predictive
31 power for clinical drug response and resistance (45). Neuroblastoma, and pediatric cancers in
32 general, are particularly amenable for genetically modelling as they are only driven by a few
33 genetic aberrations. Tumors arising in the Th-*MYCN* mouse model of neuroblastoma are

1 stroma-poor, undifferentiated or poorly differentiated with a high mitosis-karyorrhexis index,
2 closely resembling the histology of *MYCN*-amplified tumors in the >18-month age group
3 (15). The high tumor *f*BV values of ~20% reported here are consistent with the unique
4 hypervascular phenotype of childhood neuroblastoma. The median endomucin staining area
5 in the Th-*MYCN* tumors (2.3%) is consistent with the mean stained area of endothelial cell
6 marker CD31 (1.7%) in a cohort of 458 primary childhood neuroblastomas (4). Structurally,
7 the large areas of irregular sinusoidal vessels found in the hemorrhagic regions of Th-*MYCN*
8 tumors emulate the major unstable vascular patterns associated with unfavorable clinical
9 prognosis. In contrast, tumors with lower R_2^* , differentiated or not, presented a more
10 normalized vascular pattern associated with favorable prognosis factors in primary
11 neuroblastoma. Our retrospective radiological analysis of 19 cases of *MYCN*-amplified
12 neuroblastoma revealed that all tumors presented with a MRI phenotype consistent with
13 hemorrhage. Additionally, our initial experience with IS-MRI in children with
14 refractory/relapse neuroblastoma demonstrated a range of tumor median R_2^* values,
15 comparable to those measured in the Th-*MYCN* model when corrected for the difference in
16 magnetic field strength (46). Collectively these data demonstrate that the Th-*MYCN* GEM
17 model of neuroblastoma recapitulates the pathophysiology of childhood neuroblastoma and
18 provides a unique clinically-relevant and information-rich platform to identify and validate
19 imaging biomarkers of the neuroblastoma microenvironment.

20 DCE-MRI and CT-based functional imaging are commonly used to evaluate tumor perfusion
21 in the clinic. Yet their associated biomarkers are difficult to interpret as they reflect both
22 vascular perfusion and permeability. More importantly, the acquisition of robust DCE-MRI
23 data can be challenging in young children (7) and CT-based functional imaging
24 methodologies should be preferentially avoided due to the higher inherent sensitivity of
25 children to the negative effects of ionizing radiation (6). The data herein provide a strong
26 rationale for the incorporation of both IS- and SC-MRI into functional imaging-embedded
27 clinical trials for new therapeutic strategies that directly or indirectly modulate neuroblastoma
28 vascular phenotype. IS-MRI is a rapid, quantitative, safe and non-invasive R_2^* relaxometry
29 measurement that has become the reference measurement to assess both liver and cardiac iron
30 content and a primary endpoint in pediatric clinical trials in highly-transfused children
31 suffering from hematological diseases who are at risk of liver and cardiac iron overload (47).
32 The multi-gradient recalled echo MRI sequence is universally available on clinical MRI
33 scanners and IS-MRI can be easily incorporated in the neuroblastoma clinic, as demonstrated

1 herein, adding only two and half minutes to the current routine scanning session. IS-MRI has
2 been incorporated in a ancillary functional imaging study of the BEACON Neuroblastoma
3 trial. This study also demonstrates that hemorrhage detectable with IS-MRI, or the absence
4 thereof, could hold prognostic/diagnostic value. Clinically, the hemorrhagic nature of
5 undifferentiated neuroblastoma is well described, yet its routine reporting remains only
6 anecdotal (48). Its value for risk-stratification could be explored by introducing non-invasive
7 IS-MRI at the time of diagnosis. As a matter of fact, introducing susceptibility-weighted
8 imaging (SW-MRI), a variant of IS-MRI, which combines the information from both the
9 magnitude and the phase of the MR signal, would quantify hemorrhage with the added value
10 of detecting calcification (opposite phase to hemorrhage). SW-MRI can assist in the
11 differential diagnosis of neuroblastoma (frequently calcified) versus Wilms' tumors (seldom
12 calcified) as this not easily detected on conventional MRI. SC-MRI, which combines R_2^*
13 mapping with the use of USPIO particles, does not require bolus timing or complex image
14 acquisition schemes, and provides an attractive steady-state imaging alternative to DCE-MRI.
15 Recent clinical studies highlighting the safe off-label use of the USPIO preparation
16 ferumoxytol for MRI in adults and children is a promising step-forward, warranting further
17 evaluation of the approach in clinical trials (49).

18 In summary, the robust MRI-pathology cross-validation methodology used in this study
19 validates IS- and SC- MRI as robust non-invasive imaging techniques to characterize,
20 quantify and map the unique vascular phenotype of neuroblastoma and its therapeutic
21 modulation. With the central role of angiogenesis in determining and predicting the clinical
22 behavior of neuroblastoma, baseline fBV and R_2^* , have the potential to provide diagnostics
23 and prognosis information at the time of diagnosis and provide currently unavailable
24 predictive and pharmacodynamic biomarkers for anti-angiogenic therapy against
25 neuroblastoma. Lastly, this study highlights the potential role of functional MRI in delivering
26 precision medicine to children with neuroblastoma.

27

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46

Acknowledgments: AstraZeneca for the supply of cediranib.

References

1. Matthay KK, Maris JM, Schleiermacher G, Nakagawara A, Mackall CL, Diller L, *et al.* Neuroblastoma. *Nat Rev Dis Primers* **2016**;2:16078
2. Meitar D, Crawford SE, Rademaker AW, Cohn SL. Tumor angiogenesis correlates with metastatic disease, N-myc amplification, and poor outcome in human neuroblastoma. *J Clin Oncol* **1996**;14:405-14
3. Peddinti R, Zeine R, Luca D, Seshadri R, Chlenski A, Cole K, *et al.* Prominent microvascular proliferation in clinically aggressive neuroblastoma. *Clin Cancer Res* **2007**;13:3499-506
4. Tadeo I, Bueno G, Berbegall AP, Fernandez-Carrobles MM, Castel V, Garcia-Rojo M, *et al.* Vascular patterns provide therapeutic targets in aggressive neuroblastic tumors. *Oncotarget* **2016**;7:19935-47
5. Jakovljevic G, Culic S, Stepan J, Bonevski A, Seiwert S. Vascular endothelial growth factor in children with neuroblastoma: a retrospective analysis. *J Exp Clin Cancer Res* **2009**;28:143
6. Brisse HJ, McCarville MB, Granata C, Krug KB, Wootton-Gorges SL, Kanegawa K, *et al.* Guidelines for imaging and staging of neuroblastic tumors: consensus report from the International Neuroblastoma Risk Group Project. *Radiology* **2011**;261:243-57
7. Miyazaki K, Jerome NP, Collins DJ, Orton MR, d'Arcy JA, Wallace T, *et al.* Demonstration of the reproducibility of free-breathing diffusion-weighted MRI and dynamic contrast enhanced MRI in children with solid tumours: a pilot study. *Eur Radiol* **2015**;25:2641-50
8. O'Connor JP, Jayson GC. Do imaging biomarkers relate to outcome in patients treated with VEGF inhibitors? *Clin Cancer Res* **2012**;18:6588-98
9. Wu EX, Tang H, Jensen JH. Applications of ultrasmall superparamagnetic iron oxide contrast agents in the MR study of animal models. *NMR Biomed* **2004**;17:478-83
10. Robinson SP, Howe FA, Griffiths JR, Ryan AJ, Waterton JC. Susceptibility contrast magnetic resonance imaging determination of fractional tumor blood volume: a noninvasive imaging biomarker of response to the vascular disrupting agent ZD6126. *Int J Radiat Oncol Biol Phys* **2007**;69:872-9
11. Robinson SP, Boulton JKR, Vasudev NS, Reynolds AR. Monitoring the Vascular Response and Resistance to Sunitinib in Renal Cell Carcinoma In Vivo with Susceptibility Contrast MRI. *Cancer Res* **2017**;77:4127-34
12. Jamin Y, Tucker ER, Poon E, Popov S, Vaughan L, Boulton JK, *et al.* Evaluation of clinically translatable MR imaging biomarkers of therapeutic response in the TH-MYCN transgenic mouse model of neuroblastoma. *Radiology* **2013**;266:130-40
13. Vlachogiannis G, Hedayat S, Vatsiou A, Jamin Y, Fernandez-Mateos J, Khan K, *et al.* Patient-derived organoids model treatment response of metastatic gastrointestinal cancers. *Science* **2018**;359:920-6
14. O'Connor JP, Aboagye EO, Adams JE, Aerts HJ, Barrington SF, Beer AJ, *et al.* Imaging biomarker roadmap for cancer studies. *Nat Rev Clin Oncol* **2017**;14:169-86

- 1 15. Moore HC, Wood KM, Jackson MS, Lastowska MA, Hall D, Imrie H, *et al.*
2 Histological profile of tumours from MYCN transgenic mice. *J Clin Pathol*
3 **2008**;61:1098-103
- 4 16. Workman P, Aboagye EO, Balkwill F, Balmain A, Bruder G, Chaplin DJ, *et al.*
5 Guidelines for the welfare and use of animals in cancer research. *Br J Cancer*
6 **2010**;102:1555-77
- 7 17. Kilkenny C, Browne WJ, Cuthill IC, Emerson M, Altman DG. Improving bioscience
8 research reporting: the ARRIVE guidelines for reporting animal research. *PLoS Biol*
9 **2010**;8:e1000412
- 10 18. Weiss WA, Aldape K, Mohapatra G, Feuerstein BG, Bishop JM. Targeted expression
11 of MYCN causes neuroblastoma in transgenic mice. *The EMBO journal*
12 **1997**;16:2985-95
- 13 19. Dreys J, Siegert P, Medinger M, Mross K, Strecker R, Zirrgiebel U, *et al.* Phase I
14 clinical study of AZD2171, an oral vascular endothelial growth factor signaling
15 inhibitor, in patients with advanced solid tumors. *J Clin Oncol* **2007**;25:3045-54
- 16 20. Tropres I, Lamalle L, Peoc'h M, Farion R, Usson Y, Decorps M, *et al.* In vivo
17 assessment of tumoral angiogenesis. *Magn Reson Med* **2004**;51:533-41
- 18 21. Otsu N. A Threshold Selection Method from Gray-Level Histograms. *IEEE*
19 *Transactions on Systems, Man and Cybernetics* **1979**;9:62-6
- 20 22. Parzen E. On Estimation of a Probability Density Function and Mode. *Ann Math*
21 *Statist* **1962**;33:1065-76
- 22 23. Chainey S, Tompson L, Uhlig S. The Utility of Hotspot Mapping for Predicting
23 Spatial Patterns of Crime. *Security Journal* **2008**;21:4-28
- 24 24. Myronenko A, Song XB. Point Set Registration: Coherent Point Drift. *Ieee T Pattern*
25 *Anal* **2010**;32:2262-75
- 26 25. Manias KA, Gill SK, MacPherson L, Foster K, Oates A, Peet AC. Magnetic
27 resonance imaging based functional imaging in paediatric oncology. *Eur J Cancer*
28 **2017**;72:251-65
- 29 26. Daudigeos-Dubus E, Le Dret L, Bawa O, Opolon P, Vievard A, Villa I, *et al.* Dual
30 inhibition using cabozantinib overcomes HGF/MET signaling mediated resistance to
31 pan-VEGFR inhibition in orthotopic and metastatic neuroblastoma tumors. *Int J*
32 *Oncol* **2017**;50:203-11
- 33 27. Daudigeos-Dubus E, Le Dret L, Lanvers-Kaminsky C, Bawa O, Opolon P, Vievard
34 A, *et al.* Regorafenib: Antitumor Activity upon Mono and Combination Therapy in
35 Preclinical Pediatric Malignancy Models. *PLoS One* **2015**;10:e0142612
- 36 28. Calero R, Morchon E, Johnsen JI, Serrano R. Sunitinib suppress neuroblastoma
37 growth through degradation of MYCN and inhibition of angiogenesis. *PLoS One*
38 **2014**;9:e95628
- 39 29. Rossler J, Monnet Y, Farace F, Opolon P, Daudigeos-Dubus E, Bourredjem A, *et al.*
40 The selective VEGFR1-3 inhibitor axitinib (AG-013736) shows antitumor activity in
41 human neuroblastoma xenografts. *Int J Cancer* **2011**;128:2748-58
- 42 30. Cazes A, Lopez-Delisle L, Tsarovina K, Pierre-Eugene C, De Preter K, Peuchmaur
43 M, *et al.* Activated Alk triggers prolonged neurogenesis and Ret upregulation
44 providing a therapeutic target in ALK-mutated neuroblastoma. *Oncotarget*
45 **2014**;5:2688-702
- 46 31. Chantry YH, Gustafson WC, Itsara M, Persson A, Hackett CS, Grimmer M, *et al.*
47 Paracrine signaling through MYCN enhances tumor-vascular interactions in
48 neuroblastoma. *Sci Transl Med* **2012**;4:115ra3
- 49 32. Di Paolo D, Ambrogio C, Pastorino F, Brignole C, Martinengo C, Carosio R, *et al.*
50 Selective therapeutic targeting of the anaplastic lymphoma kinase with liposomal

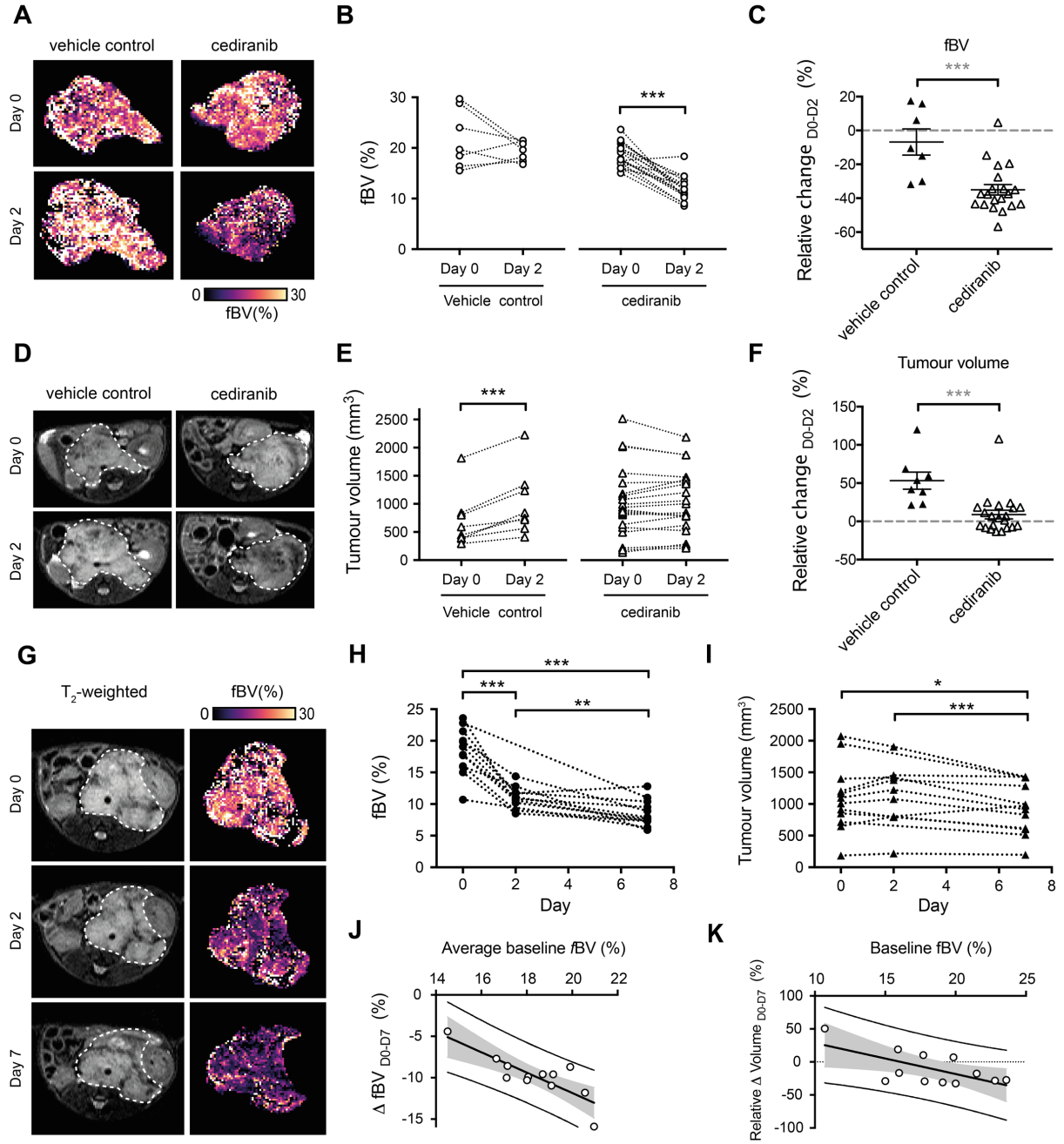
- 1 siRNA induces apoptosis and inhibits angiogenesis in neuroblastoma. *Mol Ther*
2 **2011**;19:2201-12
- 3 33. Jamin Y, Glass L, Hallsworth A, George R, Koh DM, Pearson AD, *et al.* Intrinsic
4 susceptibility MRI identifies tumors with ALKF1174L mutation in genetically-
5 engineered murine models of high-risk neuroblastoma. *PLoS One* **2014**;9:e92886
- 6 34. Moreno L, Caron H, Geoerger B, Eggert A, Schleiermacher G, Brock P, *et al.*
7 Accelerating drug development for neuroblastoma - New Drug Development
8 Strategy: an Innovative Therapies for Children with Cancer, European Network for
9 Cancer Research in Children and Adolescents and International Society of Paediatric
10 Oncology Europe Neuroblastoma project. *Expert Opin Drug Discov* **2017**;12:801-11
- 11 35. Bid HK, Kerk S. BET bromodomain inhibitor (JQ1) and tumor angiogenesis.
12 *Oncoscience* **2016**;3:316-7
- 13 36. Romain C, Paul P, Kim KW, Lee S, Qiao J, Chung DH. Targeting Aurora kinase-A
14 downregulates cell proliferation and angiogenesis in neuroblastoma. *J Pediatr Surg*
15 **2014**;49:159-65
- 16 37. Chen Z, Zhao Y, Yu Y, Pang JC, Woodfield SE, Tao L, *et al.* Small molecule
17 inhibitor regorafenib inhibits RET signaling in neuroblastoma cells and effectively
18 suppresses tumor growth in vivo. *Oncotarget* **2017**;8:104090-103
- 19 38. Kortlever RM, Sodir NM, Wilson CH, Burkhart DL, Pellegrinet L, Brown Swigart L,
20 *et al.* Myc Cooperates with Ras by Programming Inflammation and Immune
21 Suppression. *Cell* **2017**;171:1301-15 e14
- 22 39. Dejure FR, Eilers M. MYC and tumor metabolism: chicken and egg. *EMBO J*
23 **2017**;36:3409-20
- 24 40. Berry T, Luther W, Bhatnagar N, Jamin Y, Poon E, Sanda T, *et al.* The
25 ALK(F1174L) mutation potentiates the oncogenic activity of MYCN in
26 neuroblastoma. *Cancer Cell* **2012**;22:117-30
- 27 41. Lopez-Delisle L, Pierre-Eugene C, Louis-Brennetot C, Surdez D, Raynal V, Baulande
28 S, *et al.* Activated ALK signals through the ERK-ETV5-RET pathway to drive
29 neuroblastoma oncogenesis. *Oncogene* **2018**
- 30 42. Huang D, Rutkowski JL, Brodeur GM, Chou PM, Kwiatkowski JL, Babbo A, *et al.*
31 Schwann cell-conditioned medium inhibits angiogenesis. *Cancer Res* **2000**;60:5966-
32 71
- 33 43. Chlenski A, Liu S, Crawford SE, Volpert OV, DeVries GH, Evangelista A, *et al.*
34 SPARC is a key Schwannian-derived inhibitor controlling neuroblastoma tumor
35 angiogenesis. *Cancer Res* **2002**;62:7357-63
- 36 44. Poliani PL, Mitola S, Ravanini M, Ferrari-Toninelli G, D'Ippolito C, Notarangelo LD,
37 *et al.* CEACAM1/VEGF cross-talk during neuroblastic tumour differentiation. *J*
38 *Pathol* **2007**;211:541-9
- 39 45. Gengenbacher N, Singhal M, Augustin HG. Preclinical mouse solid tumour models:
40 status quo, challenges and perspectives. *Nat Rev Cancer* **2017**;17:751-65
- 41 46. Almeida GS, Panek R, Hallsworth A, Webber H, Papaevangelou E, Boulton JK, *et al.*
42 Pre-clinical imaging of transgenic mouse models of neuroblastoma using a dedicated
43 3-element solenoid coil on a clinical 3T platform. *Br J Cancer* **2017**;117:791-800
- 44 47. Verlhac S, Morel M, Bernaudin F, Bechet S, Jung C, Vasile M. Liver iron overload
45 assessment by MRI R2* relaxometry in highly transfused pediatric patients: an
46 agreement and reproducibility study. *Diagn Interv Imaging* **2015**;96:259-64
- 47 48. Peuchmaur M, d'Amore ES, Joshi VV, Hata J, Roald B, Dehner LP, *et al.* Revision of
48 the International Neuroblastoma Pathology Classification: confirmation of favorable
49 and unfavorable prognostic subsets in ganglioneuroblastoma, nodular. *Cancer*
50 **2003**;98:2274-81

- 1 49. Toth GB, Varallyay CG, Horvath A, Bashir MR, Choyke PL, Daldrup-Link HE, *et al.*
2 Current and potential imaging applications of ferumoxytol for magnetic resonance
3 imaging. *Kidney Int* **2017**;92:47-66
- 4 50. Allen LM, Hasso AN, Handwerker J, Farid H. Sequence-specific MR imaging
5 findings that are useful in dating ischemic stroke. *Radiographics* **2012**;32:1285-97;
6 discussion 97-9
7
8

1 **Figures**

2

Figure 1



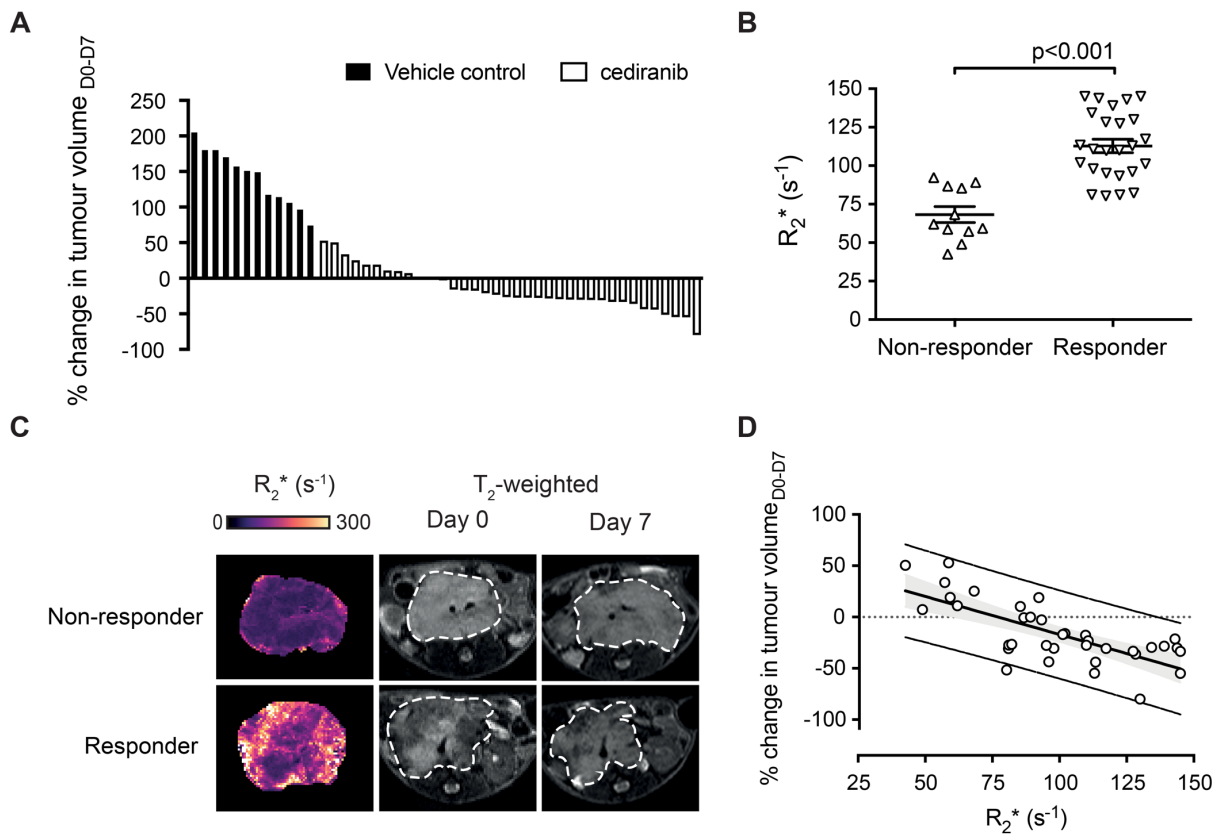
3

4

1 **Fig. 1.** Susceptibility-contrast MRI-derived fractional blood volume (*f*BV) predicts response
2 to vascular endothelial growth factor receptor (panVEGFR) inhibitor cediranib in the
3 Th-*MYCN* model of neuroblastoma. (A) Representative tumor parametric *f*BV maps
4 in the Th-*MYCN* mice prior to (Day 0) and 24h (Day 2) after treatment with 6mg/kg
5 cediranib or vehicle. Changes (B) and relative changes (C) in tumor median *f*BV
6 prior to (Day 0) and 24h (Day 2) after treatment with 6mg/kg cediranib or vehicle.
7 (D) Representative T₂-weighted anatomical MRI taken through the abdomen of
8 tumor-bearing Th-*MYCN* mice prior to (Day 0) and 24h (Day 2) after treatment with
9 6mg/kg cediranib or vehicle. (E) Changes and (F) relative changes in tumor volume
10 prior to (Day 0) and 24h (Day 2) after treatment with 6mg/kg cediranib or vehicle.
11 Data are the individual median value for each tumor and the cohort mean ± 1 s.e.m.
12 (**p<0.005, two-tailed paired Student's t-test; **p<0.01, ***p<0.005, two-tailed
13 unpaired Student's t-test, 5% level of significance). (G) Representative T₂-weighted
14 anatomical MRI and tumor *f*BV parametric map of a Th-*MYCN* mouse prior, 24h and
15 7 days after daily treatment with 6mg/kg cediranib. (H) Changes in tumor *f*BV and
16 (I) tumor volume in Th-*MYCN* mice prior, 24h and 7 days after daily treatment with
17 6mg/kg cediranib. (J) Average tumor median *f*BV correlated with change in *f*BV over
18 the 7 days of the trial ($r = -0.83$, $p = 0.0008$). (K) Tumor median *f*BV correlated with
19 change in tumor volume over the 7 days of the trial ($r = -0.65$, $p = 0.02$).
20
21

1

Figure 2



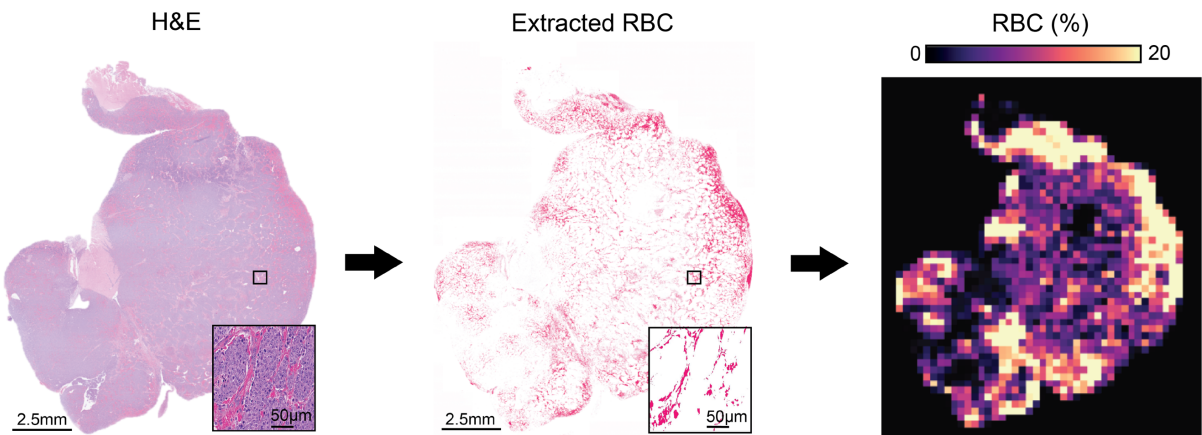
2

3

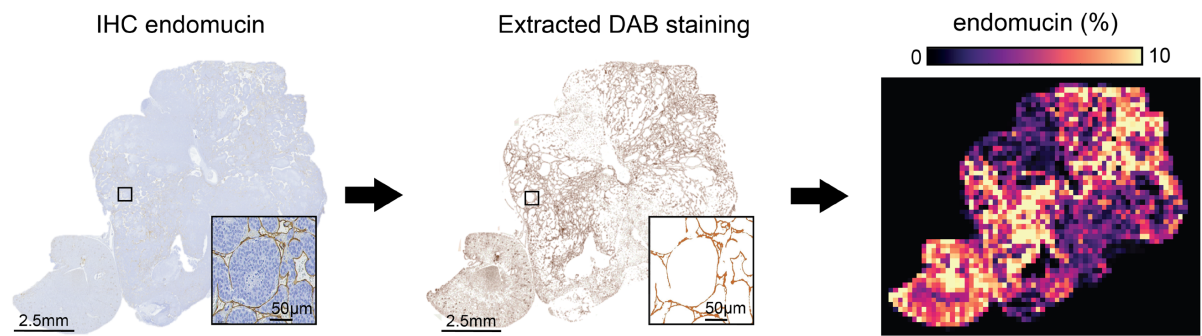
1 **Fig. 2.** Intrinsic susceptibility MRI-derived transverse relaxation rate R_2^* predicts response to
2 vascular endothelial growth factor receptor (panVEGFR) inhibitor cediranib in the
3 Th-*MYCN* model of neuroblastoma. (A) Waterfall plot documenting relative changes
4 in tumor volume in the Th-*MYCN* mouse model of neuroblastoma following seven-
5 day treatment with daily dose of 6mg/kg cediranib or vehicle ($\Delta\text{Volume}_{\text{cediranib_D0-D7}} = -$
6 $16 \pm 5\%$ vs $\Delta\text{Volume}_{\text{vehicle_D0-D7}} = 142 \pm 11\%$, $p < 0.0001$, two-tailed unpaired Student's t-
7 test with a 5% level of significance). (B) Baseline R_2^* value (Day 0) of responsive
8 and non-responsive tumors (Data are the individual median value for each tumor and
9 the cohort mean ± 1 s.e.m., $p < 0.0001$, two-tailed unpaired Student's t-test with a 5%
10 level of significance). (C) Representative baseline native R_2^* maps for responsive and
11 progressive tumors in Th-*MYCN* transgenic mice (assessed by anatomical MRI)
12 following daily treatment with 6mg/kg cediranib for 7 days. (D) Native tumor median
13 R_2^* (Day 0) correlated with relative change in tumor volume following daily
14 treatment with 6mg/kg cediranib for 7 days ($r = -0.72$, $p < 0.001$).
15

Figure 3

A



B



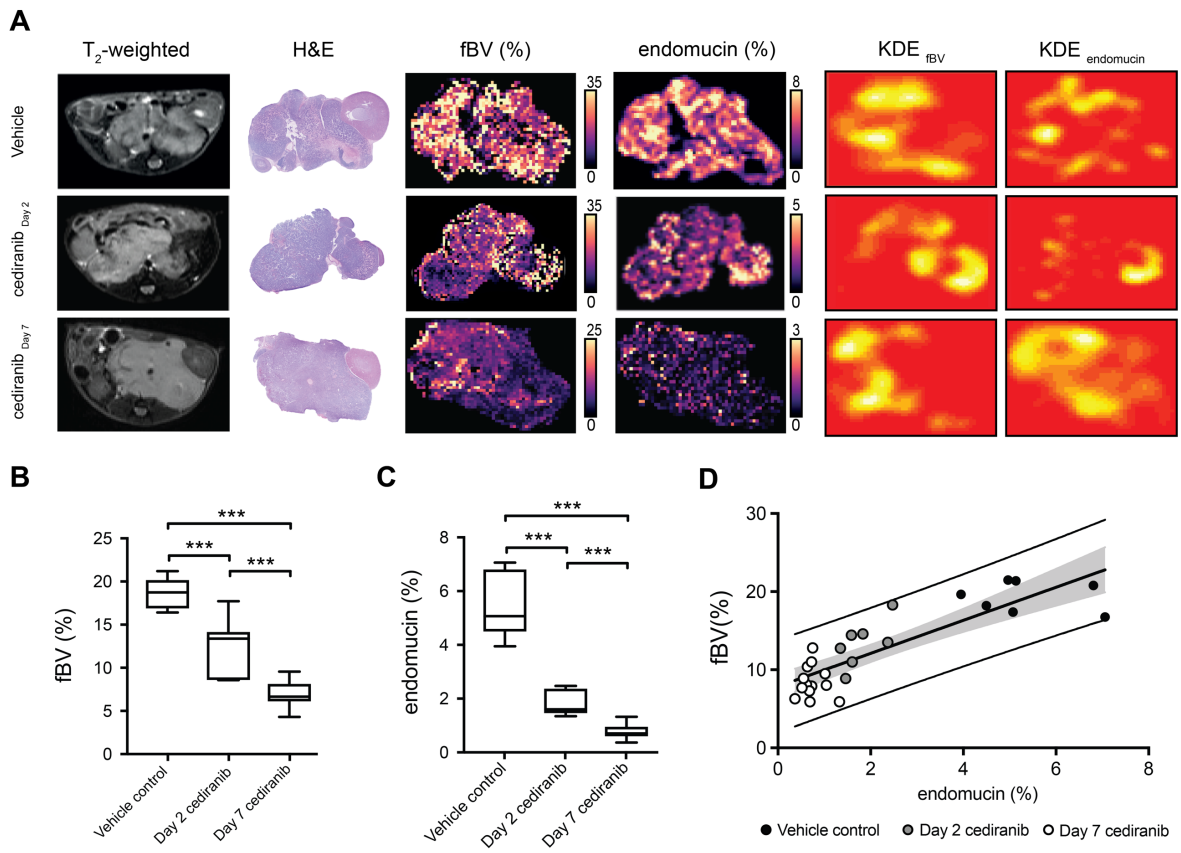
- 1
- 2
- 3
- 4

1 **Fig. 3.** Pipeline for the generation of computed maps of red blood cells (RBC) and
2 endomucin staining from histopathological images in Th-*MYCN* tumors. **(A)** RBC
3 were automatically extracted from hematoxylin and eosin (H&E)-stained whole
4 section slides with an accuracy of 99.74%. **(B)** Endomucin staining was extracted from
5 chromogenic immunohistochemistry whole section slides with an accuracy of 97%.
6 Note that the contrast of the whole slide extracted endomucin was enhanced, solely for
7 printing visibility. Whole-section images were subsequently processed to match the
8 MRI resolution, with the fraction of pixels occupied by red blood cells **(A)** and
9 endomucin **(B)** within 518x518 pixels of the original images, representing a single
10 pixel in the final maps.

11

1

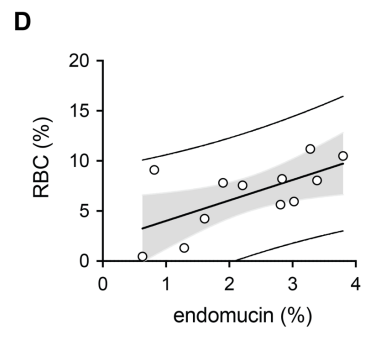
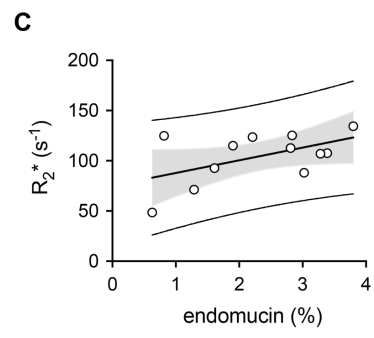
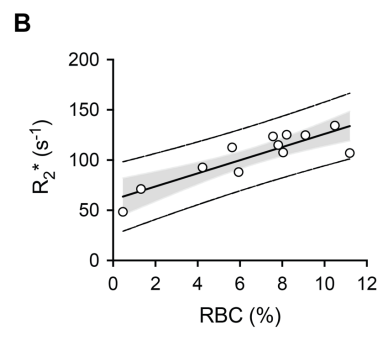
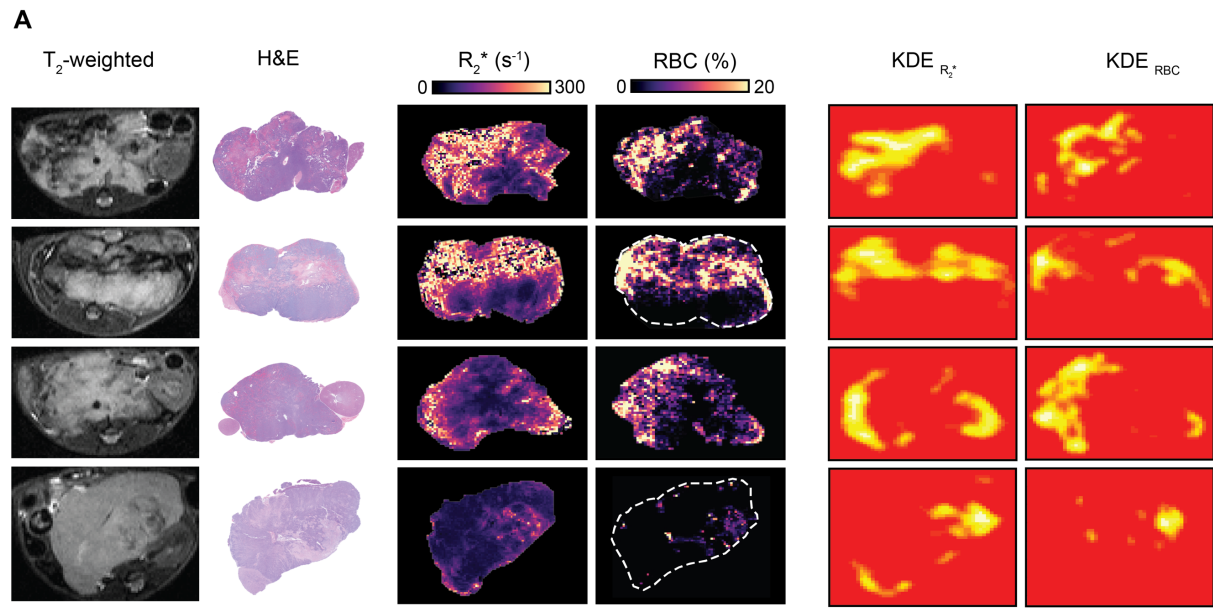
Figure 4



2

1 **Fig. 4.** Susceptibility-contrast MRI-derived fractional blood volume (fBV) correlates with
2 endothelial cell marker endomucin fraction in tumors of the Th-*MYCN* model of
3 neuroblastoma (A), Representative T₂-weighted anatomical MRI images and their
4 corresponding H&E-stained histology, parametric fBV maps and computed areas of
5 vascular endothelial cell marker endomucin staining, kernel density estimation (KDE)
6 hotspot maps of high values (above the 85th percentile of each tumor sample) of fBV
7 and endomucin fraction in the Th-*MYCN* model of neuroblastoma, 24 hours and 7
8 days after daily treatment with cediranib or vehicle (24 hours). (Note the different
9 scale for fBV and endomucin maps across the groups to better capture the
10 intratumoral heterogeneity). (B) Median tumor fBV and (C) median tumor endomucin
11 fraction area in the Th-*MYCN* GEM model 24 hours and 7 days after treatment with
12 cediranib or vehicle (***p*<0.001, two-tailed unpaired Student's t-test). (D) Tumor
13 median fBV correlated with endomucin fraction area (*r*= 0.85, *p*<0.0001).
14

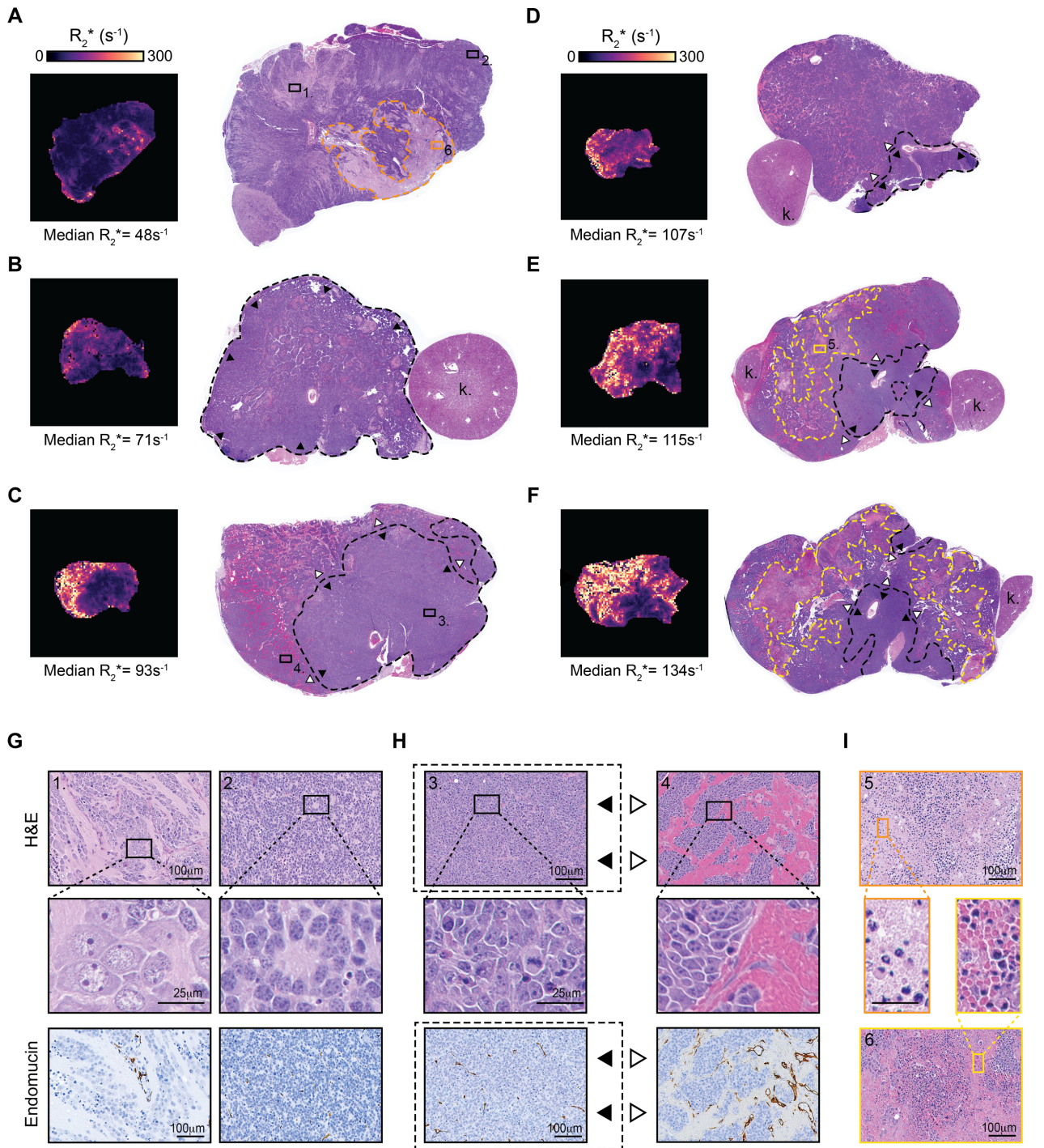
Figure 5



1
2

1 **Fig. 5.** Intrinsic susceptibility MRI-derived transverse relaxation rate R_2^* correlates with
2 tumor red blood cells content. (A) Representative T_2 -weighted MRI images and their
3 corresponding H&E histology, maps of the transverse relaxation rate R_2^* and
4 computed area of red blood cells (RBC) detected from H&E stained images, kernel
5 density estimation (KDE) hotspot maps of high values (above the 85th percentile of
6 each tumor sample) of R_2^* and RBC fraction in tumors arising in control Th-*MYCN*
7 transgenic mouse. (B) Tumor median R_2^* positively correlated with the mean value of
8 RBC fraction detected from H&E stained tumors arising in the Th-*MYCN* transgenic
9 mouse model, $r = 0.87$, $p = 0.0003$. R_2^* values (C) and mean value of RBC fraction
10 (D) also correlated with endomucin staining. ($r = 0.53$, $p = 0.08$ and $r = 0.64$, $p =$
11 0.02).
12

Figure 6



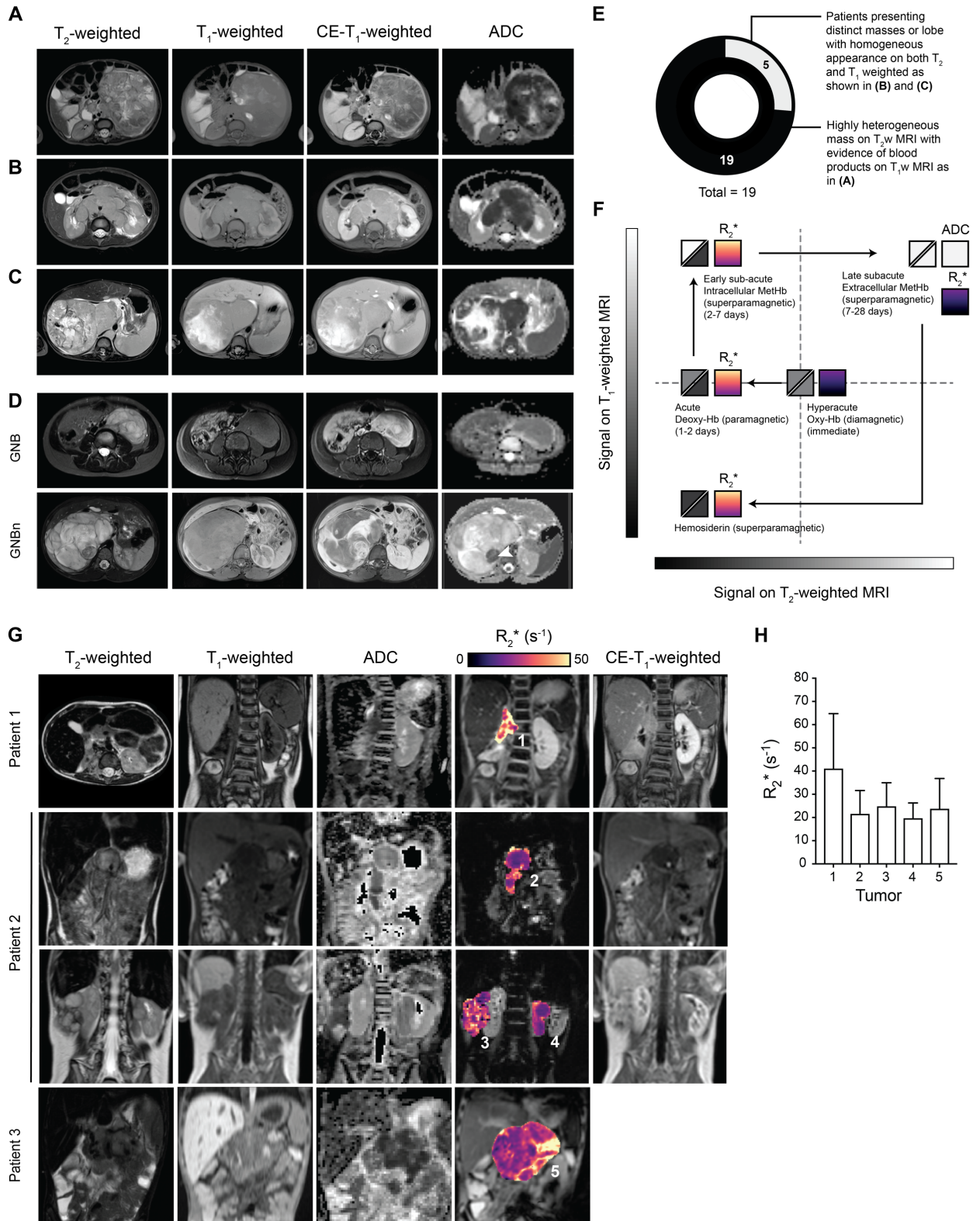
1
2

1 **Fig. 6.** Calibration of intrinsic susceptibility MRI-derived tumor R_2^* phenotype with
2 histopathological phenotype in the Th-*MYCN* genetically-engineered murine model of
3 neuroblastoma. (A-F) show susceptibility-weighted MRI-derived maps of the tumor
4 transverse relaxation R_2^* and their corresponding hematoxylin and eosin (H&E)-
5 stained whole-section slides (k., kidney), arranged by increasing median tumor R_2^*
6 value. Tumors with very low median value of R_2^* ($< 60s^{-1}$), (A) present with a
7 differentiating phenotype characterized by (G) large areas of differentiating
8 neuroblasts, with enlarged cytoplasm and more prominent nucleoli, surrounded by
9 large amounts of neuropil (1.), alongside dense regions of undifferentiated neuroblast
10 (2.). Note that hemorrhage is absent from these tumors and the vasculature is
11 characterized by sparse capillaries-like blood vessels as revealed by
12 immunohistochemistry staining for the vascular endothelial cells marker endomucin.
13 For a median value of R_2^* of $\sim 70 s^{-1}$ and above, tumors present with dense area of
14 poorly differentiated neuroblast presenting with numerous presence of mitotic
15 features (H), yet tumor median R_2^* is determined by the increasing ratio of region
16 with large area of hemorrhage and high vascular density (4., ---, \triangleright) over region with
17 lower density of capillary like vessel (3., ---, \blacktriangleright). Note that the extravasated red blood
18 cells (RBCs) appear intact. Additionally, tumors with median R_2^* over $110s^{-1}$ show
19 vast area of cellular damage (necrosis) filled with intact and damaged RBCs (I) (---,
20 5). Necrotic area with low RBC content (---, 6) only present with small regional
21 increase in R_2^* (as seen in (A)).

22

23

Figure 7



1 **Fig. 7.** Hemorrhage in *MYCN*-amplified childhood neuroblastoma MRI and initial experience
2 with SW-MRI in the neuroblastoma clinic. **(A-C)** show abdominal axial fat-
3 suppressed STIR T₂-weighted-MRI images, fat-suppressed SPAIR T₁-weighted MRI
4 images, before and after administration of gadolinium (Gd)-based contrast agent, and
5 diffusion-weighted MRI-derived apparent diffusion coefficient (ADC) maps of
6 children with neuroblastoma (NB) at the time of diagnosis. **(A)** *MYCN*-amplified
7 high-risk neuroblastoma in a 10-month old boy, **(B)** a 2-year old boy, **(C)** a 8-month-
8 old boy, **(D)** Comparison between a ganglioneuroblastoma (GNB) in a 4-year-old
9 female patient and ganglioneuroblastoma nodular (GNBn) in a 5-year-old male patient.
10 Note the presence of the neuroblastic nodule in the GNBn easily identified on ADC
11 maps (arrow head) **(E)** Proportion of patients with *MYCN*-amplified neuroblastoma
12 (n=19) presenting with a MRI phenotype suggestive of the presence or absence of a
13 hemorrhagic phenotype, based on the well-established knowledge of the appearance
14 of ageing blood in hematoma on conventional MRI (50) as illustrated in **(F)**. **(G)**
15 show abdominal T₂-weighted and T₁-weighted images (not fat-suppressed), ADC
16 maps, intrinsic susceptibility MRI-derived transverse relaxation R₂* map of tumors,
17 and post Gd contrast-enhanced T₁-weighted MRI images in children with
18 refractory/relapsing neuroblastoma. (Patient 1: 7-year old male, Patient 2: 5-year old
19 male, Patient 3: 6-year old female). **(H)** Tumor median R₂* values for each individual
20 tumor shown in **(G)** (\pm SD). Note that R₂* values increased monotonically and
21 approximately linearly with magnetic field strength-dependent and as such clinical
22 R₂* value at 1.5T are estimated to be four times lower that if they were measured at
23 7T (46).

24