

## 1 **S - Material and Methods**

### 2 *v) DNA Extraction*

3 DNA was extracted from either 2 mm punches taken from tissue in paraffin blocks with an Integra  
4 miltex biopsy punch (Integra, 33-31-P/25), marked areas from unstained paraffin sections mounted on  
5 glass slides (USS) or from fresh frozen tissue. Tissue from paraffin punches was deparaffinized by  
6 immersion in 100% xylene twice for 30 minutes and in 100% ethanol twice for 30 min, and air dried.  
7 The tissue punches were then incubated 1- 3 days at 56°C in 50-200  $\mu$ l (volume depending on tissue  
8 block size) of proteinase K digestion buffer (PKDB) composed with 1x PCR buffer II (Applied  
9 Biosystems 4486220), 0.05% NP-40 and 200 $\mu$ g/ml proteinase K (Qiagen 19133). After incubation the  
10 proteinase was heat inactivated for 10 min at 96°C. The sample was let to cool to room temperature  
11 and then centrifuged for 1 min at 13.000 rpm. Tissue sections on USS were deparaffinized on slides,  
12 aligned to a corresponding slide stained with hematoxylin and eosin to demarcate tumour and normal  
13 adjacent regions and then scraped off into 30-100  $\mu$ l PKDB buffer and treated the same as above.  
14 Fresh frozen tissue was added directly from frozen to PKDB buffer and processed similarly.

15

### 16 *vi) RNA Extraction from Fresh Frozen Tissue*

17 RNA from fresh frozen (FF) tissue was isolated with the Direct-zol RNA miniprep Kit (R2050, Zymo  
18 Research). A 25-50 mg block of FF tissue was transferred to a 50 ml Corex centrifuge tube with  
19 600  $\mu$ l of Biorad PureZol RNA Isolation reagent (Cat#732-6890) and homogenised with a small  
20 homogeniser tip for 3-4 x 20 sec burst with intermittent 30 sec cool down periods on ice. The tube  
21 was centrifuged for 5 min at 4°C at 3000 rpm and the supernatant transferred to a tube containing  
22 600  $\mu$ l 100% ethanol and mixed by inversion. The mix was transferred to a Zymo-Spin IIC Column  
23 and centrifuged for 1 min at 8000 rpm. After one wash with 400  $\mu$ l RNA wash buffer a DNaseI  
24 treatment was performed (5  $\mu$ l of DNaseI +75  $\mu$ l of DNase buffer/column) for 15 min at RT. One  
25 wash with 400  $\mu$ l RNA pre-wash buffer, one wash with 700  $\mu$ l RNA wash buffer with a 2 min  
26 centrifugation was performed before elution with 100 $\mu$ l RNase free water. Eluted RNA was

27 quantified by Nanodrop ND-1000 Spectrophotometer and Quality assessed with RNA Tapescreen<sup>(TM)</sup>  
28 on a TapeStation<sup>(TM)</sup>. For RNA extraction FFPE tissue scraped from slides (USS) or punched from  
29 paraffin blocks (as described above) were processed with the RNAsort kit from CellDataSci  
30 (CD501) according to the manufacturer's protocol.

31

32 *vii) Bisulfite conversion*

33 200 ng DNA was bisulfite modified with either the Qiagen Epitect Bisulfite kit (Cat 59104) or the  
34 Zymo Research EZ DNA Methylation kit (D5001) according to the manufacturers' instructions.  
35 Complete bisulfite modification was monitored by an internal bisulfite control position in the  
36 pyrosequencing assays for MGMT and SDHC.

37

38 *viii) Analysis of MGMT and SDHC promoter methylation*

39 *MGMT*: A 124 bp sized PCR amplicon located in the promoter region of the MGMT gene was  
40 amplified from 1-25 ng bisulfite converted DNA with 375 nM forward primer, with a 20-mer 5' M13  
41 overhang to facilitate post PCR sequencing,  
42 (TGTAACGACGGCCAGTTTATAGTTTTYGGATATGTTGGGATAG) and 187.5nM of  
43 biotinylated reverse primer ([btn]-TCCCAAACACTCACCAAATC) with the Qiagen PyroMark kit  
44 (Qiagen 978703).

45 *SDHC*: For methylation analysis of the SDHC gene a 134 bp PCR amplicon located in the promoter  
46 region of the SDHC gene was amplified from 2-25 ng of bisulfite converted DNA with 375 nM  
47 forward primer, with a 20-mer 5' M13 overhang to facilitate post PCR sequencing,  
48 (TGTAACGACGGCCAGTTTATAGGAGAAGTTTTAGAGTTTTTTAAAGAG) and 250nM of  
49 biotinylated reverse primer ([btn]-AAAATAACRCCAAACRACCCC). The PCR conditions were  
50 7 min at 95°C, followed by 20 sec at 95°C, 30 sec at 53°C, and 20 sec at 72°C for 40 cycles, and an  
51 end incubation at 72°C for 5 min. The resulting PCR amplicon was quality assessed for purity and

52 yield on a 2% agarose gel. For MGMT, a nested sequencing primer  
53 (GTTTTTAGAACGTTTTGYGTTT) was used to analyse 4 CpGs in 10  $\mu$ l of the PCR sample on the  
54 Qiagen Q24 pyrosequencer (sequence to analyse: YGAYGTTYGTAGGTTTTYGT). For SDHC a  
55 nested sequencing primer (GTTATATGATATTTTAAATTT) was used to analyse six CpGs  
56 (sequence to analyse: YGATTTTTAGTYGGYGYGTTTTYGTTTYGGGT). Fully methylated and  
57 unmethylated human control DNA that had been treated with bisulfite were used as controls on each  
58 pyrosequencing run. Pyrosequencing results were confirmed in a subset of cases by Sanger  
59 sequencing with the M13 forward primer recognising the M13 extension of the forward oligos for  
60 both the MGMT and the SDHC PCR amplicon.

61

62 *ix) MGMT expression analysis with quantitative RT-PCR*

63 RNA (125-250 ng) was transcribed into cDNA with random hexamers employing the High Capacity  
64 cDNA Reverse Transcription Kit from Applied Biosystems (ref 4368814) according to manufacturer's  
65 instructions. cDNA was diluted tenfold after synthesis and 1% was used in each RT-PCR well.  
66 Relative MGMT expression was analyzed according to (Uno et al 2011) (16) with SYBR Green using  
67 the PowerUp SYBR Green Master Mix (Applied Biosystems, ref 01061935), with MGMT oligo's;  
68 forward 5'-GCTGAATGCCTATTTCCACCA-3'/reverse 5'-CACAACTTCAGCAGCTTCCA-3';  
69 normalised to the average Ct value of three internal reference genes; hypoxanthine guanine  
70 phosphoribosyltransferase (HPRT1); forward 5'- GAAAAGGACCCACGAAGTGT -3'/ reverse 5'-  
71 AGTCAAGGGCATATCCTACAACA -3', beta-glucuronidase (GUSB); forward 5'-  
72 AAAATACGTGGTTGGAGAGCTCATT -3'/ reverse 5'- CCGAGTGAAGATCCCCTTTTTA -3'and  
73 TATA-box binding protein (TBP); forward 5'- AGGATAAGAGAGCCACGAACCA -3'/ reverse 5'-  
74 CTTGCTGCCAGTCTGGACTGT -3'. The delta Ct was calculated by subtracting the mean of  
75 triplicate Ct values for MGMT with the mean of the triplicate Ct values of all 3 reference genes.  
76 MGMT promoter hypermethylation was defined as a mean MGMT methylation across CpG islands 1-  
77 8 of > 10% for this study.

78

79 *Clinical germline DNA sequencing*

80 DNA was extracted from peripheral blood samples according to standard protocols. Next generation  
81 sequencing of a clinical gene panel including; *SDHA*, *SDHB*, *SDHC*, *SDHD*, *KIT*, *PDGFRA* and *NF1*  
82 was performed by the laboratory staff at Cambridge University Hospital NHS Foundation Trust or  
83 Birmingham Women's and Children's Hospital NHS Trust using the TrusightOne or Trusight Cancer  
84 sequencing panels (Illumina Inc., UK). An average coverage depth of >20 fold was achieved for 98%  
85 of the regions sequenced. All detected variants were confirmed by Sanger sequencing. Whole exon  
86 deletions and duplications and large rearrangements are not detected using this method and multiple  
87 ligation probe analysis (MLPA) was performed for *SDHB*, *SDHC* and *SDHD*.

88 *xii) SDHB Immunohistochemistry*

89 SDHB immunohistochemistry (IHC) was performed on 4 µm sections of FFPE tissue using a  
90 commercially available SDHB polyclonal rabbit antibody (Sigma-Aldrich HPA002868). Heat-  
91 induced epitope retrieval was carried out using a Leica heat retrieval solution (HPA002867, Sigma  
92 Aldrich, UK). SDHB deficiency was defined by a loss or abnormal IHC staining pattern for SDHB in  
93 the tumour cells when compared to the staining pattern in tumour vasculature cells.

94