Supplementary Table 2 – Next Generation Sequencing (NGS) result of MMR-deficient cases (excluding cases with MLH1 promoter hypermethylation)

Variants are either likely pathogenic (class 4) or pathogenic (class 5) unless otherwise specified.

Abbreviations: VAF = Variant allele frequency. LOH = Loss of heterozygosity. SNP = Single Nucleotide Polymorphism. NP = not performed. VAF = Variant allele freque Immunohistochemistry results: + = normal nuclear staining. - = loss of staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls. +/++ = weak staining in neoplastic cells with positive internal controls.

Study ID	Resection or biopsy	Immu	nohisto	chemistry pa	ittern		NGS results neoplastic tissue	Variant detected in			
		PMS2	MLH1	MSH6	MSH2	Gene	Variant	VAF : coverage	LOH	non-neoplastic tissue	
3	Resection	+	np	+/++	-	MSH2	Nonsense variant*	0.779:715	Probable based on 1 SNP and VAF	Yes	
18	Biopsy	-	-	+	np		No relevant variants detected				
22	Resection	+	nn	_	+/++	MSH2	NM_000251.2:c.1777C>T	0.480:125	No	No	
33			np	-	+/++	1013112	Deletion exon 1			No	
46	Resection	-	+	+	np		NGS data of insufficient quality				
48	Resection	-	+	+	+	MLH1	NM_000249.3:c.112A>C	0.48:448	No informative SNPs	Normal tissue not available	
71	Resection	+	np	-	+/++		No relevant variants detected				
	Resection	Failed	Failed np			MLH1	Missense variant classified as pathogenic by InSiGHT	0.479:1308	No informative SNPs	Yes	
85				np -	+		NM_000249.3:c.1513_1520dup	0.168:1985		No	
						MSH6	C-deletion				
94	Resection	-	-	+	np	MLH1	NM_000249.3:c.676C>T		yes	Not performed	
00	Resection	h +	+ np			+/++	MSH6	Frameshift variant*	0.483:1989	N	Yes
98				- H	+/++	INISHO	NM_000179.2:c.3743del	0.329:1989	No	No	
118	Resection	+	np	-	-		NGS data of insufficient quality				
110	Resection		_	_	nn	PMS2	Frameshift variant*	0.498:1933	yes	Yes	

113	nesection		T	, T	IIΡ	MSH2	NM_000251.2:c.187dup	0.204:1967		No
124	Resection	+	np	-	failed		No relevant variants detected			According to PA-report this is a Lynch
156	Resection	+	np	-	-	MSH2	NM_000251.2:c.2027C>G	0.219:283	No informative SNPs	Not performed
206	Resection	-	-	+	np	MLH1	Frameshift variant*	0.429:1919	No informative SNPs	Yes
211	Resection	-	-	+	np	MLH1	NM_000249.3:c.454-13A>G	0.918:244	No informative SNPs, VAF is however	Normal tissue not available
214	Resection	subclo nal -	+	+	np	MSH2	NM_000251.2:c.1414C>T (class 3 VUS)	0.156:257		Not performed
226	Resection	+	nn		_	MSH6	Frameshift variant*	0.511:1621		Yes
230			np	-	-	IVISHO	NM_000179.2:c.3172G>T (class 3 VUS)	0.313:1995		No
240	Resection	-	+		200	DMC2	NM_000535.5:c.2287G>T	0.159:1233	No informative SNPs	Normal tissue not
249				+	np	PMS2	NM_000535.5:c.1882C>T	0.397:315		available
316	Resection	+	np	+/++	-	MSH2	Exon deletion*	Not applicable	Yes	Yes
325	Resection	+	np	-	-		No relevant variants detected			
333	Resection	+	np	-	-	MSH6	NM_000179.2:c.3128del	0.185:352	No informative SNPs	Not performed
335	Resection	+	np	-	-	MSH2	Missense variant classified as likely pathogenic by InSiGHT*	0.520:1997	LOH probable based on 1 informative SNP	Yes
344	Resection	-	-	+	np	MLH1	NM_000249.3:c.94_110del	0.341:1510	No informative SNPs	Not performed
363	Resection	+	np	-	-	MSH2	Frameshift variant*	0.499:914	No based on 1 SNP	Yes
270	Resection	-	- +	+	np	PMS2	Nonsense variant*	0.500:1225	No	Yes
379							NM_000535.5:c.1802C>G	0.421:680		No
414	Resection		nn		4/11	MSH6	Frameshift variant*	0.481:1795	No bosed on 1 CAID	Yes
414		+	np		+/++	IVISHO	NM_000179.2:c.3533del	0.239:1980	No based on 1 SNP	No

426	Resection	-	-	+	np	MLH1	Frameshift variant*	0.539:1990	No informative SNPs	Yes						
453	Resection	-	-	+	np	MLH1	NM_000249.3:c.791-2A>C (class 3 VUS)	0.634:1994	No informative SNPs	No						
460	Resection	+	np	+/++	-	MSH2	NM_000251.2:c.2557G>T	0.241:1312	Unlikely based on 1 SNP	Not performed						
466	Resection	-	-	+	np		No relevant variants detected									
474	Resection	+	np	-	-	MSH2	Frameshift variant*	0.691:676	Probable based on 1 SNP	Yes						
490	Posaction			Subclonal	np	MLH1	Nonsense variant*	0.744:1999	yes	Yes						
460	Resection			-	пр	MSH6	No relevant variants detected		yes							
526	Resection	+	np	-	-		No relevant variants detected									
551	Resection	-	-	+	Np	MLH1	NM_000249.3:c.2145_2168del	0.578:211	No informative SNPs	Normal tissue not available						
558	Resection	-	+	Subclonal -	+/++	PMS2	NM_000535.5:c.638del	0.378:1995	Possibly based on 3 SNPs	No						
568	Resection	-	+/++	+	np	PMS2	NM_000251.2:c.2458+1G>A	0.381:1998	No	Not performed						
FOF	Resection	+	22		+/++	MSH6	Frameshift variant*	0.453:1190	- No	Yes						
393			np	-		IVISHO	NM_000179.2:c.2232G>T (class 3 VUS)	0.169:349		No						
FOC	Resection			np	-		NM_000251.2:c.1861C>T 0.491:1611	Nor	Normal tissue not							
590		+	+			-	MSH2	NM_000251.2:c.2458+1G>A	0.271:399	No	available					
504	Resection	+	np									MSH2	NM_000251.2:c.1601G>A	0.346:619	No informative SNPs	Not a sufferment
601				-	-	MSH6	NM_000179.2:c.1436_1440del	0.35:1980	Yes	Not performed						
687	Resection	-	np	-	-		NGS data of insufficient quality									
698	Resection	+	np	Subclonal	+		NGS data of insufficient quality									
710	Resection	-	+	+	np	PMS2	NM_000535.5:c.1405A>T	0.522:1994	No	Normal tissue not available						

720 Resection	+	np	-	-	MSH2	Nonsense variant*	0.678:1772	No	Yes
722 Resection		np	_	+/++	MSH6	Frameshift variant*	0.532:342	no	Yes
722 Resection	+		-	+/++	IVISHO	NM_000179.2:c.1444C>T	0.285:895		No
723 Resection	-	-	+	np	MLH1	Frameshift variant*	0.937:1449	Probable based on one SNP and VAF of	Yes
746 Resection	-	-	+	np	MLH1	Frameshift variant*	0.594:1721	Yes	Yes
748 Resection	+	np	-	-		NGS data of insufficient quality			
760 Resection	-	-	+	np	MLH1	NM_000249.3:c.252del	0.498:601	yes	No