

Department of Pathology and Laboratory Medicine No.201, Sec. 2, Shipai Rd., Beitou District, Taipei City, Taiwan 11217, R.O.C.

Tel: 02-2875-7449

Date: 03 Dec 2021 1 of 3

Sample Information

Patient Name: 傅仙婷 Gender: Female ID No.: A224622912 History No.: 37771929

Age: 43

Ordering Doctor: DOC2589K 曾令民

Ordering REQ.: D6H4D9G Signing in Date: 2021/12/03

Path No.: S110-94701 **MP No.:** BR21053

Assay: Oncomine BRCA1/2 Assay

Sample Type: Blood

Date of blood drawing: 2021/11/25

Reporting Doctor: DOC5452C 周德盈 (Phone: 8#5452)

Note:

Sample Cancer Type: Breast Cancer

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Relevant Biomarkers

No clinically significant biomarkers found in this sample.

Variant Details

DNA Sequence Variants Allele **Amino Acid Change** Variant Effect ClinVar1 Gene Coding Locus Frequency Transcript Coverage BRCA2 p.(?) c.-26G>A chr13:32890572 49.61% NM_000059.3 unknown Benign 3995 BRCA2 p.(N372H) c.1114A>C chr13:32906729 Benign 3998 48.65% NM 000059.3 missense BRCA2 p.(K1132=) c.3396A>G chr13:32911888 51.60% NM_000059.3 synonymous Benign 3990 BRCA2 p.(L1521=) c.4563A>G chr13:32913055 99.88% NM_000059.3 Benign 3474 synonymous BRCA2 p.(P1842L) c.5525C>T chr13:32914017 48.89% NM_000059.3 missense Uncertain 3768 significance

¹ Based on Clinvar version 20200329

Variant Details (continued)

DNA Sequence Variants (continued)

				Allele				
Gene	Amino Acid Change	Coding	Locus	Frequency	Transcript	Variant Effect	Clinvar	Coverage
BRCA2	p.(V2109I)	c.6325G>A	chr13:32914817	49.16%	NM_000059.3	missense	Conflicting interpretations of pathogenicity	3983
BRCA2	p.(V2171=)	c.6513G>C	chr13:32915005	99.93%	NM_000059.3	synonymous	Benign	3998
BRCA2	p.(S2414=)	c.7242A>G	chr13:32929232	53.86%	NM_000059.3	synonymous	Benign	3999
BRCA2	p.(V2466A)	c.7397T>C	chr13:32929387	99.82%	NM_000059.3	missense	Benign	3997
BRCA2	p.(P3292L)	c.9875C>T	chr13:32972525	48.12%	NM_000059.3	missense	Benign	3969
BRCA1	p.(S1613G)	c.4837A>G	chr17:41223094	50.73%	NM_007294.4	missense	Benign	3992
BRCA1	p.(S1436=)	c.4308T>C	chr17:41234470	50.27%	NM_007294.4	synonymous	Benign	4000
BRCA1	p.(K1183R)	c.3548A>G	chr17:41244000	50.48%	NM_007294.4	missense	Benign	3998
BRCA1	p.(E1038G)	c.3113A>G	chr17:41244435	50.21%	NM_007294.4	missense	Benign	3999
BRCA1	p.(P871L)	c.2612C>T	chr17:41244936	48.85%	NM_007294.4	missense	Benign	3998
BRCA1	p.(L771=)	c.2311T>C	chr17:41245237	50.65%	NM_007294.4	synonymous	Benign	3996
BRCA1	p.(S694=)	c.2082C>T	chr17:41245466	49.81%	NM_007294.4	synonymous	Benign	3997

¹ Based on Clinvar version 20200329

Date: 03 Dec 2021

Signatures

Testing Personnel:

Laboratory Supervisor:

Pathologist: