P. O. Box 570 Jefferson City, MO 65102-0570 (573) 522-2845

ENROLLMENT SITE/SATELLITE SITE (NAME AND ADDRESS)	REFERRING PROVIDER (FOR DIRECT BILLING)				
A. PERSONAL DATA					
NAME (LAST, FIRST, MIDDLE INITIAL)	SOCIAL SECURITY NUMBER				
DATE OF BIRTH MM DD YYYY CLIENT ELIGIBILITY VERIFIED INSURANCE COVERAGE DEDUCTIBLE MET Yes No Yes No	REFERRAL FEE MEDICARE				
VISITTYPE ☐ Initial ☐ Annual ☐ Rescreen ☐ Navigation only Height Weight Bloc	od Pressure 1st Reading/ Average /				
B. BREAST CANCER SCREENING					
B 1. Does client report any BSE symptoms? Yes No (If "YES" complete B2.)					
B 2. Symptoms Reported By Client (Check any that apply. If 1, 2, 3 or 4B is checked, may have two (2) diagnost	tics at clinician's discretion.)				
□ (1) Lump □ (4A) Pain/Tenderness - 1st occurrence □ (4B) Pain/Tenderness - 2nd occurrence					
☐ (2) Nipple discharge ☐ (5) Other (specify) ☐ (3) Skin changes (dimpling, retraction, new nipple inversion,					
ulceration, Paget's disease)					
B 3. CBE within normal limits and findings Present at CBE (check yes or no and one explanation) Date of CBE/(MM/DD/YYYY)					
□ Yes □ Within normal limits					
□(1) Benign finding (fibrocystic changes, diffuse lumpiness, clearly defined thickening, tenderness or nodularity) □ No - Suspicious for cancer (Any checked findings requires completion of two (2) diagnostic procedures entered on purple breast form.)					
□ (2) Discrete palpable mass (includes masses that may be diffuse, poorly defined thickening, cystic or solid)					
iower tran usuar, prominent ver	ins, unilateral; unusual increase in size, unilateral palpable supraclavicular, infraclavicular or axillary				
☐ (4) Nipple or areolar scaliness or erythema lymph nodes; also swelling of u	_ (v)g,,,				
☐ Focal pain and tenderness Rescreen CRF Planned ☐ Yes ☐ No / Diagnostic Workup Planned ☐	Voc. □ No. /				
Rescreen CBE Planned □ Yes □ No/ Diagnostic Workup Planned □ (must be less than 10 months) MM YYYY (must be less than 10 months)	MM YYYY				
B 4. High Risk for Breast Cancer ☐ (1) Yes* ☐ (2) No ☐ (9) Not assessed/Unknown * At least one must be met: BRCA Mutation, First Degree Releative BRCA Carrier, or Greater Than 20-26 Percent Lifetime Risk					
B 5. Mammogram					
Previous mammogram					
Type of mammogram ☐ Screening ☐ Diagnostic ☐ Tomosynthesis Method used for mammogr	ram □ Digital □ Conventional				
Mammography provider facility	□ Mammogram Van				
(facility name / city)					
□ (4) Mammogram not done or CBE done and □ (5) Cervical record only, no breast service provided diagnostic workup planned □ (6) Referred to direct biller					
(1) Routine screening mammogram (3) Abnormal mammogram done by a non-program funded provider, patient referred in for					
☐ (2) Mammogram performed to evaluate symptoms: ☐ Personal history of breast cancer ☐ Previous abnormal mammogram results (rescreen) ☐ Date client referred for diagnosis//					
	MM DD YYYY				
SMHW mammogram result (check one) (results with * require additional follow-up) Reporting Only					
Left Right (Indicate why only one breast had mammogram in COMMENTS) Left Right Normal □ □ (1) Negative (Category 1) Abnormal □ □ (3) Probably Benign (Category 3)					
□ □ (2) Benign Finding (Category 2) □ □ (4) Suspicious Abnormality (Category 4)*					
	estive of Malignancy (Category 5)* y-not interpreted, repeat (Not Paid)				
	tion or film comparison (<i>Category 0</i>)				
Rescreen mammogram planned					
(must be less than 10 months) (must be less than 60 days) /	L R				
Referred for diagnostic testing/direct bill	☐ ☐ (2) Berlight finding (category 2) ☐ ☐ (3) Probably Benign (Category 3)				
(physician / facility name) ☐ MRI (High Risk ONLY. Prior authorization required.)/ Report results here	□□ (4) Suspicious (Category 4) □□ (5) Highly Suspicious (Category 5) □□ (6) Known Malignancy (Category 6)				
MRI Type: ☐ Unilateral ☐ Not Done MM DD YYYY ☐ Bilateral	(0) Kilowii Maiigrialicy (Category 6) (7) Incomplete (Category 0) (8) Results Pending				
i Dilateral	□ □ (9) Not Done				

MO 580-1799 (11-22)

□ (6) Breast and Pelvic exam only (1) Routine Pap test (screening) (No Cervical Service) □ (2) Patient under surveillance for previous abnormal (rescreen) (5) Pap test not done. Patient proceeded directly for diagnostic work-up or HPV testing			High Risk for Cervical Cancer		
— ·	Pap after primary HPV positive	,	· ·	☐ (1) Yes ☐ (2) No	
□ (3) Non-program Pap referred in f	or diagnostic evaluation	/ /	(2) Not assessed/	
□ (9) Unknown		MM DD YYYY	unknown	
C 1. Pelvic Exam Results		C 2. Pelvic Exam Findings		☐ Reporting Only	
Pelvic Exam WNL? ☐ Yes ☐ (Additional information required in "No" sel		Findings Present at Pelvic Exam (check or 1) Cervix	nly one)		
Hysterectomy? □ Yes □	l No	□ a) Polyp □] f) Ectropion		
☐ Cervix absent		☐ b) Leukoplakia (white lesions) ☐			
☐ Cervix absent due to cervical cancer		□ c) Friable □			
(needs annual Pap test)		☐ d) Ulceration ☐			
☐ Cervix present		□ e) Exophytic growth			
☐ Reason for hysterectomy unknow	vn				
Date of Pelvic Exam /	1	☐ 2) Exam Complicated by Obesity			
	DD YYYY	Rescreen planned ☐ Yes ☐ N	0 /		
Reproductive Status (check one)		. 100 _ 11	MM YYYY		
□ a) Premenopausal		Diagnostic planned ☐ Yes ☐ N	0 / /		
☐ b) Postmenopausal		(must be less than 60 days)	MM DD YYYY		
C 3. Pap Test Results				☐ Reporting Only	
Previous Pap test ☐ Yes ☐] No □ Unknown	Date of last Pap test/	Date of this Pap test		
		MM YYYY		MM DD YYYY	
Specimen adequacy	orv	Specimen type ☐ Conven	tional Smear		
	ctory due to	☐ Liquid B	ased		
□ Unknown	otory due to				
□ Annual Pap due to previous treatment for cervical cancer					
Pap test result (check one) (Results wi	th (*) require additional follow-u	p)			
Normal □ (1) Negativ	ve for intraepithelial lesion or ma	alianancy			
. , ,	natory/Infection/Reactive Change	9 .			
. ,	I Squamous Cells of Undetermi		Squamous Cell Cancer*		
(ASC-US)(May have HPV test) □ (8) Atypical Glandular Cells* (including atypicalendocer					
☐ (4) Lowgr	ade SIL (HPV/Mild Dysplasia/C	IN I)*	adenocarcinoma in situ and ade	enocarcinoma)	
☐ (5) Atypica	al Squamous Cells, cannot ex	CIUGE HSIL (ASC-H)"	Adenocarcinoma in situ*		
☐ (6) Highgr	ade SIL (with features suspicion	us for invasion/CIN II-III/CIS)*	Adenocarcinoma*		
Foderanded Calls		□ (11)	Other		
Endocervical Cells	,				
C 4. HPV Test Date	/MM/DD/YYYY			☐ Reporting Only	
Indication for HPV Test □ (1) Cotesting	/Screening HPV Test Resu	ılt	HPV DNA Genotype 16 or 18		
☐ (2) Reflex		with genotyping	negative and HPV High Risk 0 ☐ Yes	roup Positive)	
☐ (3) Not Done		0 0 11 0			
☐ (9) Unknown	(z) Negative	= (0) Olimilowii	□ No		
	genotyp	•	☐ No Test Performe	d 	
Rescreen Pap planned	□ No/_ MM YYYY	Diagnostic work-up planned (must be less than 60 days)	□ Yes □ No .	MM DD YYYY	
Referred for diagnostic work-up/ (physician/facility name)	direct biller				
Date of next routine Pap screening	/	_			

MO 580-1799 (11-22)

D. COMMENTS